

(19)



Europäisches Patentamt

European Patent Office

Office européen des brevets



(11)

EP 1 179 594 A1

(12)

EUROPEAN PATENT APPLICATION

published in accordance with Art. 158(3) EPC

(43) Date of publication:

13.02.2002 Bulletin 2002/07

(51) Int Cl.7: **C12N 15/45, C12P 21/02**

(21) Application number: **00929790.4**

(86) International application number:

PCT/JP00/03194

(22) Date of filing: **18.05.2000**

(87) International publication number:

WO 00/70055 (23.11.2000 Gazette 2000/47)

(84) Designated Contracting States:

**AT BE CH CY DE DK ES FI FR GB GR IE IT LI LU
MC NL PT SE**

Designated Extension States:

AL LT LV MK RO SI

(30) Priority: **18.05.1999 JP 20074099**

(71) Applicant: **Dnavec Research Inc.**

Ibaraki 305-0856 (JP)

(72) Inventors:

• **KITAZATO, Kaio DNAVEC Research Inc.**
Tsukuba-shi Ibaraki 305-0856 (JP)

• **SHU, Tsugumine DNAVEC Research Inc.**
Tsukuba-shi Ibaraki 305-0856 (JP)

• **KUMA, Hidekazu**

A 203, Hisamitsu Pharma. Co., Inc.
Ryugasaki-shi Ibaraki 301-0044 (JP)

• **UEDA, Yasuji**

Tsukuba-shi Ibaraki 305-0856 (JP)

• **ASAKAWA, Makoto**

Toyonaka-shi Osaka 561-0825 (JP)

• **HASEGAWA, Mamoru DNAVEC Research Inc.**

Tsukuba-shi Ibaraki 305-0856 (JP)

• **IIDA, Akihiro DNAVEC Research Inc.**

Tsukuba-shi Ibaraki 305-0856 (JP)

• **HIRATA, Takahiro DNAVEC Research Inc.**

Tsukuba-shi Ibaraki 305-0856 (JP)

(74) Representative: **VOSSIUS & PARTNER**

Siebertstrasse 4

81675 München (DE)

(54) **RNP ORIGINATING IN PARAMYXOVIRUS**

(57) A functional RNP containing negative-strand single-stranded RNA derived from Sendai virus, which has been modified so as not to express any envelope protein, has been successfully prepared. An RNP com-

prising a foreign gene is prepared and inserted into a cell with the use of a cationic liposome, thereby successfully expressing the foreign gene.

EP 1 179 594 A1

DescriptionTechnical Field

5 **[0001]** The present invention relates to paramyxovirus-derived ribonucleoprotein complex and the utilization thereof.

Background Art

10 **[0002]** Paramyxovirus is a virus comprising negative-strand RNA as the genome. Negative-strand RNA viral vectors have several characteristics significantly different from retroviruses, DNA viruses or positive-strand RNA virus vectors. Genomes or antigenomes of negative-strand RNA viruses do not directly function as mRNA, so they cannot initiate the synthesis of viral proteins and genome replication. Both RNA genome and antigenome of these viruses always exist in the form of a ribonucleoprotein complex (RNP), so they hardly cause problems caused by antisense strands, such as interfering with the assembly of genome to RNP due to mRNA hybridizing with naked genomic RNA, as in the case of positive strand RNA viruses. These viruses comprise their own RNA polymerases, performing the transcription of viral mRNAs or replication of viral genomes using RNP complex as the template. Worthy of mentioning is that negative-strand RNA (nsRNA) viruses proliferate only in the cytoplasm of host cells, causing no integration thereof into chromosomes, because they do not go through a DNA phase. Furthermore, no homologous recombination among RNAs has been recognized. These properties are considered to contribute a great deal to the stability and safety of negative-strand RNA viruses as gene expressing vectors.

20 **[0003]** Among negative-strand RNA viruses, present inventors have been focusing their attention on the Sendai virus (SeV). Sendai virus is a non-segmented type negative-strand RNA virus belonging to the genus Paramyxovirus, and is a type of murine parainfluenza virus. This virus has been said to be non-pathogenic towards humans. In addition, an attenuated laboratory strain (Z strain) of Sendai virus has been isolated, which only induces mild pneumonia in rodents, the natural hosts (J. General Virology (1997) 78, 3207-3215). This strain has been widely used as a research model for molecular level studies of the transcription-replication mechanism of paramyxoviruses. Sendai virus attaches to the host cell membrane and cause membrane fusion via its two envelope glycoproteins, hemagglutinin-neuraminidase (HN) and fusion protein (F), and efficiently releases its own RNA polymerase and RNA genome existing in the form of ribonucleoprotein complex (RNP) into the cytoplasm of host cells to carry out transcription of viral mRNA and genome replication therein (Bitzer, M. et. al., J. Virol. 71(7): 5481-5486, 1997).

30 **[0004]** Present inventors have hitherto developed a method for recovering infectious Sendai virus particles from cDNA corresponding to Sendai virus genome. In this method, for example, after infecting LLC-MK2 cells with vaccinia virus encoding T7 RNA polymerase, the cells are further transfected simultaneously with three plasmids encoding the antigenome of Sendai virus under the control of T7 promoter, the nucleoprotein (NP) of Sendai virus and the RNA polymerase proteins (P and L), respectively, to form antigenomic ribonucleoprotein complexes (RNPS) as intermediates of viral genome replication in the cells, and then replicate biologically active (functional) genomic RNPs capable of initiating viral protein transcription and virus particle assembly. When recovering the wild-type Sendai virus, these functional genomic RNPs are injected into chorioallantoic sac of chicken eggs together with reconstituted cells to perform virion multiplication (Kato, A. et al., Genes cells 1, 569-579 (1996)).

40 **[0005]** However, Sendai virus has been known to incorporate host cell proteins thereto during virion formation (J. Biol. Chem. (1997) 272, 16578-16584), and such incorporated proteins maybe possible causes of antigenicity and cytotoxicity when transferred to target cells.

45 **[0006]** In this regard, in spite of the obvious need existing for the use of RNP as vectors without utilizing Sendai virus particles, there has been no report on such a utilization.

Disclosure of the Invention

50 **[0007]** An objective of the present invention is to isolate an RNP deriving from a virus belonging to the family Paramyxoviridae, and to provide the utilization thereof as a vector. In a preferred embodiment, vectors comprising a complex of RNP with a cationic compound are provided.

[0008] The present inventors have prepared RNPs from Sendai virus belonging to the family Paramyxoviridae and investigated their use as a vector.

55 **[0009]** Specifically, first, the present inventors prepared a Sendai virus genomic cDNA deficient in the gene for the F protein, which is one of the envelope proteins of the virus, so as not to produce wild-type Sendai viruses in target cells, and further constructed a vector to express the cDNA in cells (GFP gene is inserted into the vector as a reporter at the F gene-deficient site). The vector thus prepared was transferred to cells expressing proteins required for RNP formation to produce an RNP comprising an F gene-deficient genome. Then, the RNP was removed from the cells by repeating cycles of freezing and thawing of the cells, mixed with a cationic lipofection reagent, and transferred to F

gene-expressing cells. As a result, the expression of GFP as a reporter was detected in the cells to which RNP was transfected.

[0010] Namely, present inventors succeeded not only in preparing functional RNP from Sendai virus, but also found a possibility to express a foreign gene comprised in RNP, even when this RNP is transferred to target cells utilizing, for example, a gene transfer reagent such as a cationic liposome, in stead of just infecting the RNP to cells as a constituting element of Sendai virus, and thus accomplished this invention.

[0011] Namely, this invention relates to paramyxovirus-derived RNP and the utilization thereof as a vector, more specifically to:

(1) a complex comprising (a) a negative-strand single-stranded RNA derived from a paramyxovirus, wherein said RNA is modified so as not express at least one of the envelope proteins of Paramyxoviridae viruses, and (b) a protein encoded by and binding to said negative-strand single-stranded RNA;

(2) a complex according to (1), wherein said negative-strand single-stranded RNA is modified so as to express NP, P and L proteins, but not F HN or M proteins, or any combination thereof;

(3) a complex according to (1) or (2), wherein said negative-strand single stranded RNA derives from the Sendai virus;

(4) a complex according to any of (1) through (3), wherein said negative strand single-stranded RNA further encodes a foreign gene;

(5) a composition for gene transfer, comprising a complex according to (4) and a cationic lipid;

(6) a composition for gene transfer, comprising a complex according to (4) and a cationic polymer; and,

(7) a method for expressing a foreign gene in a cell, comprising the step of introducing the composition for gene transfer according to (5) or (6) into a cell.

[0012] "NP, P, M, F, HN and L genes" of viruses belonging to the family Paramyxoviridae refer to genes encoding nucleocapsid, phospho, matrix, fusion, hemagglutinin-neuraminidase and large proteins, respectively. Respective genes of viruses belonging to subfamilies of the family Paramyxoviridae are represented in general as follows. NP gene is generally described also as the "N gene".

Genus Respirovirus	N	P/C/V	M	F	HN	-	L
Genus Rubellavirus	N	P/V	M	F	HN	(SH)	L
Genus Morbillivirus	N	P/C/V	M	F	H	-	L

[0013] Database accession numbers for nucleotide sequences of genes of the Sendai virus classified into Respirovirus of the family Paramyxoviridae are, M29343, M30202, M30203, M30204, M51331, M55565, M69046 and X17218 for NP gene, M30202, M30203, M30204, M55565, M69046, X00583, X17007 and X17008 for P gene, D11446, K02742, M30202, M30203, M30204, M69046, U31956, X00584 and X53056 for M gene, D00152, D11446, D17334, D17335, M30202, M30203, M30204, M69046, X00152 and X02131 for F gene, D26475, M12397, M30202, M30203, M30204, M69046, X00586, X02808 and X56131 for HN gene, and D00053, M30202, M30203, M30204, M69040, X00587 and X58886 for L gene.

[0014] The present invention relates to a ribonucleoprotein complex (RNP) derived from viruses belonging to the family Paramyxoviridae deficient in any of the envelope genes. The complex is modified so as not to produce the virus having the envelope protein in target cells in the absence of the envelope protein. That is, RNP according to this invention comprises (a) a negative-strand single-stranded RNA originating in paramyxovirus modified so as not to express at least one of envelope proteins of Paramyxoviridae viruses (b) a protein encoded by and binding to said negative-strand single-stranded RNA.

[0015] Proteins capable of binding to a negative-strand single-stranded RNA refer to proteins binding directly and/or indirectly to the negative-strand single-stranded RNA to form an RNP complex with the negative-strand single-stranded RNA. In general, negative-strand single-stranded RNA (genomic RNA) of paramyxovirus is bound to NP, P and L proteins. RNA contained in this RNP serves as the template for transcription and replication of RNA (Lamb, R. A., and D. Kolakofsky, 1996, *Paramyxoviridae*: The viruses and their replication, pp. 1177-1204. In *Fields Virology*, 3rd edn. Fields, B. N., D. M. Knipe, and P. M. Howley et al. (ed.), Raven Press, New York, N. Y.). Complexes of this invention include those comprising negative-strand single-stranded RNAs originating in paramyxovirus and proteins also originating in paramyxovirus which bind to the RNAs. Complexes of this invention are RNP complexes comprising, for example, negative-strand single-stranded RNA to which these proteins (NP, P and L proteins) are bound. In general,

RNP complexes of paramyxovirus are capable of autonomously self-replicating in host cells. Thus, RNPs transferred to cells intracellularly proliferate to increase the copy number of the gene (RNA contained in RNP complex), thereby leading to a high level expression of a foreign gene from RNP carrying the foreign gene. Vectors of this invention are preferably those capable of replicating RNA comprised in complexes (RNP) in transfected cells.

[0016] The origin of RNP complexes of this invention is not limited as long as it is a virus of family Paramyxoviridae, but Sendai virus belonging to the genus Paramyxovirus is especially preferred. Besides Sendai virus, RNPs of this invention may derive from the measles virus, simian parainfluenza virus (SV5) and human parainfluenza virus type 3, but the origin is not limited thereto.

[0017] Negative-strand single-stranded RNAs contained in RNPs of this invention are constructed so as to suppress the expression of at least one of the envelope proteins of Paramyxoviridae viruses. Examples of envelope proteins the expressions of which are suppressed are, F protein, HN protein, or M protein, or any combination thereof. negative-strand single-stranded RNAs are constructed so as to express NP, P and L proteins that are required for the formation of RNPs. negative-strand single-stranded RNAs contained in RNPs of this invention may be modified, for example, so as to express NP, P and L proteins, but not F and/or HN proteins.

[0018] In the case of Sendai virus (SeV), the genome of the natural virus is approximately 15,000 nucleotides in size, and the negative-strand comprises six genes encoding NP (nucleocapsid), P (phospho), M (matrix), F (fusion), HN (hemagglutinin-neuraminidase) and L (large) proteins lined in a row following the 3'-short leader region, and a short 5'-trailer region on the other end. In this invention, this genome can be modified so as not to express envelope proteins by designing a genome deficient in any of F, HN and M genes, or any combination thereof. Deficiency in either F gene or HN gene, or both is preferred. Since these proteins are unnecessary for the formation of RNP, RNPs of this invention can be manufactured by transcribing this genomic RNA (either positive or negative-strand) in the presence of NP, P and L proteins. RNP formation can be performed, for example, in LLC-MK2 cells, or the like. NP, P and L proteins can be supplied by introducing to cells expression vectors carrying the respective genes for these proteins (cf. examples). Each gene may be also incorporated into chromosomes of host cells. NP, P and L genes to be expressed for the formation of RNP need not be completely identical to those genes encoded in the genome contained in RNP. That is, amino acid sequences of proteins encoded by these genes may not be identical to those of proteins encoded by RNP genome, as long as they can bind to the genomic RNA and are capable of replicating RNP in cells, and may have mutations or may be replaced with a homologous gene from other viruses. Once an RNP is formed, NP, P and L genes are expressed from this RNP to autonomously replicate RNP in the cells.

[0019] To reconstitute and amplify an RNP in cells, the RNP is either transferred to cells (helper cells) expressing envelope proteins whose expression is suppressed by modifying negative-strand single-stranded RNA contained in the RNP, or the RNP can be reconstituted in these cells. For example, to amplify RNP from negative-strand single-stranded RNA which has been modified so as not to express F gene, F protein is arranged to be expressed together with NP, P and L proteins in the cells. Thus, a viral vector retaining envelope proteins is constructed, and amplified via its infection to helper cells.

[0020] In addition, it is also possible to use envelope proteins different from that whose expression was suppressed by modifying negative-strand single-stranded RNA. There is no particular limitation on the type of such envelope proteins. One example of other viral envelope proteins is the G protein (VSV-G) of vesicular stomatitis virus (VSV). RNP complexes of this invention can be amplified, for example, using cells expressing the G protein (VSV-G) of VSV.

[0021] Complexes of this invention can be usually prepared by (a) introducing a vector DNA encoding paramyxovirus-derived negative-strand single-stranded RNA that has been modified so as not to express at least one of the viral envelope proteins of Paramyxoviridae viruses, or a complementary strand of said RNA, into cells (helper cells) expressing envelope proteins to express the RNAs, and (b) culturing the cells to recover RNP complexes from the culture supernatant or cell extracts. By coexpressing NP, L and P proteins at the time of vector DNA expression, RNPs are formed and a virus having envelope proteins is constructed.

[0022] Vector DNA to be expressed in helper cells encodes negative-strand single-stranded RNA (negative-strand) or complementary strand thereof (positive strand) contained in complexes of this invention. Although the strand to be transcribed inside cells may be either positive or negative-strand, it is preferable to arrange so as to transcribe the positive strand for the improvement of complex reconstitution efficiency. For example, DNA encoding negative-strand single-stranded RNA or complementary strand thereof is linked downstream of T7 promoter to be transcribed to RNA by T7 RNA polymerase.

[0023] For example, a virus comprising RNP complex can be reconstituted by transfecting a plasmid expressing a recombinant Sendai virus genome deficient in envelope genes into host cells, together with a vector expressing the deficient envelope protein and NP, P/C and L protein expression vectors. Alternatively, RNP complex can be manufactured using, for example, host cells incorporated with F gene into chromosomes thereof. Amino acid sequences of these protein groups supplied from outside the viral genome need not be identical to those deriving from the virus. As long as these proteins are equally active to or more active than natural type proteins in transferring nucleic acids into cells, genes encoding these proteins may be modified by introducing a mutation or replacing with homologous genes

from other viruses. Since, in general, it has been known that long-term culture of host cells is sometimes difficult because of cytotoxicity and cell shape-altering activity of envelope proteins, they may be arranged to be expressed only when the vector is reconstituted under the control of an inducible promoter (cf. examples).

[0024] Once RNP or virus comprising RNP is formed, complexes of this invention can be amplified by introducing this RNP or virus again into the aforementioned helper cells and culturing them. This process comprises the steps of (a) introducing either the complex of this invention or viral vector comprising the complex to cells expressing envelope proteins, and (b) culturing the cells and recovering virus particles from the culture supernatant or cell extracts.

[0025] RNP may be introduced to cells as a complex formed together with, for example, lipofectamine and polycationic liposome. Specifically, a variety of transfection reagents can be utilized. Examples thereof are DOTMA (Boehringer), Superfect (QIAGEN #301305), DOTAP, DOPE, DOSPER (Boehringer #1811169), etc. Chloroquine may be added to prevent RNP from decomposition in endosomes (Calos, M. P., 1983, Proc. Natl. Acad. Sci. USA 80: 3015).

[0026] Once a viral vector is thus constructed in host cells, complexes of this invention or viral vector comprising the complexes can be further amplified by coculturing these cells with cells expressing envelope proteins. As described in Example 12, a preferable example is the method of overlaying cells expressing envelope proteins on virus producing cells.

[0027] Complexes of this invention, for example, may comprise a viral gene encoded in RNA in the complex that has been modified to reduce the antigenicity or enhance the RNA transcription and replication efficiency.

[0028] Complexes of this invention may include RNA encoding a foreign gene in their negative-strand single-stranded RNA. Any gene desired to be expressed in target cells may be used as the foreign gene. For example, when gene therapy is intended, a gene for treating an objective disease is inserted into the vector DNA encoding RNA contained in complexes. In the case where a foreign gene is inserted into the vector DNA, for example, Sendai virus vector DNA, it is preferable, to insert a sequence comprising a nucleotide number of a multiple of six between the transcription termination sequence (E) and transcription initiation sequence (S), etc. (Journal of Virology, Vol. 67, No. 8, 1993, p. 4822-4830). The foreign gene may be inserted before or after each of the viral genes (NP, P, M, F, HN and L genes) (cf. examples). E-I-S sequence (transcription initiation sequence-intervening sequence-transcription termination sequence) or portion thereof is appropriately inserted before or after a foreign gene so as not to interfere with the expression of genes before or after the foreign gene. Expression level of the inserted foreign gene can be regulated by the type of transcription initiation sequence added upstream of the foreign gene, as well as the site of gene insertion and nucleotide sequences before and after the gene. For example, in Sendai virus, the nearer the insertion site is to the 3'-end of negative-strand RNA (in the gene arrangement on the wild type viral genome, the nearer to NP gene), the higher the expression level of the inserted gene is. To secure a high expression level of a foreign gene, it is preferable to insert the foreign gene upstream of NP gene (the 3'-side in negative-strand) or between NP and P genes. Conversely, the nearer the insertion position is to the 5'-end of negative-strand RNA (in the gene arrangement on the wild type viral genome, the nearer to L gene), the lower the expression level of the inserted gene is. To suppress the expression of a foreign gene to a low level, the foreign gene is inserted, for example, to the far most 5'-side of the negative-strand, that is, downstream of L gene in the wild type viral genome (the 5'-side adjacent to L gene in negative-strand) or upstream of L gene (the 3'-side adjacent to L gene in negative-strand). To facilitate the insertion of a foreign gene, a cloning site may be designed at the inserting position. Cloning site can be arranged to be, for example, the recognition sequence for restriction enzymes. Foreign gene fragments can be inserted into the restriction enzyme site in the vector DNA encoding the genome. Cloning site may be arranged to be a so-called multi-cloning site comprising a plurality of restriction enzyme recognition sequences. RNA genome in complexes of this invention may harbor at the insertion sites foreign genes other than those described above.

[0029] Viral vectors comprising RNP complex derived from recombinant Sendai virus carrying a foreign gene can be constructed as follows according to, for example, the description in "Kato, A. et al., 1997, EMBO J. 16: 578-587" and "Yu, D. et al., 1997, Genes Cells 2: 457-466".

[0030] First, a DNA sample comprising the cDNA nucleotide sequence of a desired foreign gene is prepared. It is preferable that the DNA sample can be electrophoretically identified as a single plasmid at concentrations of 25 ng/μl or more. Below, a case where a foreign gene is inserted to DNA encoding viral genome utilizing NotI site will be described as an example. When NotI recognition site is included in the objective cDNA nucleotide sequence, it is preferable to delete the NotI site beforehand by modifying the nucleotide sequence using site-specific mutagenesis and such method so as not to alter the amino acid sequence encoded by the cDNA. From this DNA sample, the desired gene fragment is amplified and recovered by PCR. To have NotI sites on the both ends of amplified DNA fragment and further add a copy of transcription termination sequence (E), intervening sequence (I) and transcription initiation sequence (S) (EIS sequence) of Sendai virus to one end, a forward side synthetic DNA sequence and reverse side synthetic DNA sequence (antisense strand) are prepared as a pair of primers containing NotI restriction enzyme cleavage site sequence, transcription termination sequence (E), intervening sequence (I), transcription initiation sequence (S) and a partial sequence of the objective gene.

[0031] For example, to secure cleavage by NotI, the forward side synthetic DNA sequence is arranged in a form in

which any two or more nucleotides (preferably 4 nucleotides excluding GCG and GCC, sequences originating in NotI recognition site, more preferably ACTT) are selected on the 5'-side of the synthetic DNA, NotI recognition site "gcg-gccgc" is added to its 3'-side, and to the 3'-side thereof, any desired 9 nucleotides or nucleotides of 9 plus a multiple of 6 nucleotides are added as the spacer sequence, and to the 3'-side thereof, about 25 nucleotide-equivalent ORF including the initiation codon ATG of the desired cDNA is added. It is preferable to select about 25 nucleotides from the desired cDNA as the forward side synthetic DNA sequence so as to have G or C as the final nucleotide on its 3'-end.

[0032] In the reverse side synthetic DNA sequence, any two or more nucleotides (preferably 4 nucleotides excluding GCG and GCC, sequences originating in the NotI recognition site, more preferably ACTT) are selected from the 5'-side of the synthetic DNA, NotI recognition site "gcggccgc" is added to its 3'-side, and to its further 3'-side, an oligo DNA is added as the insertion fragment to adjust the length. This oligo DNA is designed so that the total nucleotide number including the NotI recognition site "gcggccgc", complementary sequence of cDNA and EIS nucleotide sequence of Sendai virus genome originating in the virus described below becomes a multiple of six (so-called "rule of six"; Kolakofski, D. et al., J. Virol. 72: 891-899, 1998). Further to the 3'-side of inserted fragment, a sequence complementary to S sequence of Sendai virus, preferably 5'-CTTTCACCCT-3', I sequence, preferably 5'-AAG-3', and a sequence complementary to E sequence, preferably 5'-TTTTTCTTACTACGG-3', is added, and further to the 3'-side thereof, about 25 nucleotide-equivalent complementary sequence counted in the reverse direction from the termination codon of the desired cDNA sequence the length of which is adjusted to have G or C as the final nucleotide, is selected and added as the 3'-end of the reverse side synthetic DNA.

[0033] PCR can be done according to the usual method with, for example, ExTaq polymerase (Takara Shuzo). Preferably, PCR is performed using Vent polymerase (NEB), and desired fragments thus amplified are digested with NotI, then inserted to NotI site of the plasmid vector pBluescript. Nucleotide sequences of PCR products thus obtained are confirmed with a sequencer to select a plasmid having the right sequence. The inserted fragment is excised from the plasmid using NotI, and cloned to the NotI site of the plasmid carrying the genomic cDNA deficient in envelope genes. Alternatively, it is also possible to obtain the recombinant Sendai virus cDNA by directly inserting the fragment to the NotI site without the mediation of the plasmid vector pBluescript.

[0034] It is also possible to transcribe a vector DNA encoding the virus genome in test tubes or cells, reconstitute RNP with viral L, P and NP proteins, and produce the virus vector comprising this RNP. Reconstitution of virus from the vector DNA can be carried out according to methods known in the art using cells expressing envelope proteins (WO97/16539 and 97/16538: Durbin, A. P. et al., 1997, Virology 235: 323-332; Whelan, S. P. et al., 1995, Proc. Natl. Acad. Sci. USA 92: 8388-3392; Schnell, M. J. et al., 1994, EMBO J. 13: 4195-4203; Radecke, F. et al., 1995, EMBO J. 14: 5773-5784; Lawson, N. D. et al., Proc. Natl. Acad. Sci. USA 92: 4477-4481; Garcin, D. et al., 1995, EMBO J. 14: 6087-6094; Kato, A. et al., 1996, Genes Cells 1: 569-579; Baron, M. D. and Barrett, T., 1997, J. Virol. 71: 1265-1271; Bridgen, A. and Elliott, R. M., 1996, Proc. Natl. Acad. Sci. USA 93: 15400-15404). When a viral vector DNA is made deficient in F, HN and/or M genes, infectious virus particles are not formed with such a defective vector. However, it is possible to form infectious virus particles and amplify the virus comprising the complex by separately transferring these deficient genes, genes encoding other viral envelope proteins and such to host cells and expressing them therein.

[0035] Methods for transferring vector DNA into cells include the following: 1) the method of preparing DNA precipitates that can be taken up by objective cells; 2) the method of preparing a DNA comprising complex which is suitable for being taken up by objective cells and which is also not very cytotoxic and has a positive charge, and 3) the method of instantaneously boring on the objective cellular membrane pores wide enough to allow DNA molecules to pass through by electric pulse.

[0036] In Method 2), a variety of transfection reagents can be utilized, examples being DOTMA (Boehringer), Superfect (QIAGEN #301305), DOTAP, DOPE, DOSPER (Boehringer #1811169), etc. An example of Method 1) is a transfection method using calcium phosphate, in which DNA that entered cells are incorporated into phagosomes, and a sufficient amount is incorporated into the nuclei as well (Graham, F. L. and Van Der Eb, J., 1973, Virology 52: 456; Wigler, M. and Silverstein, S., 1977, Cell 11: 223). Chen and Okayama have investigated the optimization of the transfer technique, reporting that optimal DNA precipitates can be obtained under the conditions where 1) cells are incubated with DNA in an atmosphere of 2 to 4% CO₂ at 35°C for 15 to 24 h, 2) cyclic DNA with a higher precipitate-forming activity than when linear DNA is used, and 3) DNA concentration in the precipitate mixture is 20 to 30 µg/ml (Chen, C. and Okayama, H., 1987, Mol. Cell. Biol. 7: 2745). Method 2) is suitable for a transient transfection. An old method is known in the art in which a DEAE-dextran (Sigma #D-9885, M.W. 5 x 10⁵) mixture is prepared in a desired DNA concentration ratio to perform the transfection. Since most of the complexes are decomposed inside endosomes, chloroquine may be added to enhance transfection effects (Calos, M. P., 1983, Proc. Natl. Acad. Sci. USA 80: 3015). Method 3) is referred to as electroporation, and is more versatile compared to methods 1) and 2) because it doesn't have cell selectivity. Method 3) is said to be efficient under optimal conditions for pulse electric current duration, pulse shape, electric field potency (gap between electrodes, voltage), conductivity of buffers, DNA concentration, and cell density.

[0037] Among the above-described three categories, transfection reagents (method 2)) are suitable in this invention,

because method 2) is easily operable, and facilitates the examining of many test samples using a large amount of cells. Preferably, Superfect Transfection Reagent (QIAGEN, Cat. No. 301305) or DOSPER Liposomal Transfection Reagent (Boehringer Mannheim, Cat. No. 1811169) is used.

[0038] Specifically, the reconstitution of the viral vector from cDNA can be performed as follows.

[0039] Simian kidney-derived LLC-MK2 cells are cultured in 24-well to 6-well plastic culture plates or 100 mm diameter culture dish using a minimum essential medium (MEM) containing 10% fetal calf serum (FCS) and antibiotics (100 units/ml penicillin G and 100 µg/ml streptomycin) to 70 to 80% confluency, and infected, for example, with recombinant vaccinia virus VTF7-3 expressing T7 polymerase at 2 PFU/cell. This virus has been inactivated by a UV irradiation treatment for 20 min in the presence of 1 µg/ml psoralen (Fuerst, T. R. et al., Proc. Natl. Acad. Sci. USA 83: 8122-8126, 1986; Kato, A. et al., Genes Cells 1: 569-579, 1996). Amount of psoralen added and UV irradiation time can be appropriately adjusted. One hour after the infection, the cells are transfected with 2 to 60 µg, more preferably 3 to 5 µg, of the above-described recombinant Sendai virus cDNA by the lipofection method and such using plasmids (24 to 0.5 µg of pGEM-N, 12 to 0.25 µg of pGEM-P and 24 to 0.5 µg of pGEM-L, more preferably 1 µg of pGEM-N, 0.5 µg of pGEM-P and 1 µg of pGEM-L) (Kato, A. et al., Genes Cells 1: 569-579, 1996) expressing trans-acting viral proteins required for the production of full-length Sendai viral genome together with Superfect (QIAGEN). The transfected cells are cultured in a serum-free MEM containing 100 µg/ml each of rifampicin (Sigma) and cytosine arabinoside (AraC) if desired, more preferably only containing 40 µg/ml of cytosine arabinoside (AraC) (Sigma), and concentrations of reagents are set at optima so as to minimize cytotoxicity due to the vaccinia virus and maximize the recovery rate of the virus (Kato, A. et al., 1996, Genes Cells 1, 569-579). After culturing for about 48 to 72 h following the transfection, the cells are recovered, disrupted by repeating three cycles of freezing and thawing, transfected to LLC-MK2 cells expressing envelope proteins, and cultured. After culturing the cells for 3 to 7 days, the culture solution is collected. Alternatively, infectious virus vectors can be obtained more efficiently by transfecting LLC-MK2 cells already expressing envelope proteins with plasmids expressing NP, L and P proteins, or transfecting together with an envelope-expressing plasmid. Viral vectors can be amplified by culturing these cells overlaid on LLC-MK2 cells expressing envelope proteins (cf. examples). Virus titer contained in the culture supernatant can be determined by measuring the hemagglutination activity (HA), which can be assayed by "endo-point dilution method" (Kato, A. et al., 1996, Genes Cells 1, 569-579). Virus stock thus obtained can be stored at -80°C.

[0040] The type of host cells used for virus reconstitution is not particularly limited, so long as RNP complex or viral vector can be reconstituted therein. For example, in the reconstitution of Sendai virus vector or RNP complex, culture cells such as simian kidney-derived CV-1 cells and LLC-MK2 cells, hamster kidney-derived BHK cells, and so on can be used. Infectious virus particles having the envelope can be also obtained by expressing appropriate envelope proteins in these cells. To obtain Sendai virus vector in a large quantity, the virus can be amplified, for example, by inoculating RNP or virus vector obtained from the above-described host cells into embryonated chicken eggs together with vectors expressing envelope genes. Alternatively, viral vectors can be produced using transgenic chicken eggs incorporated with envelope protein genes. Methods for manufacturing viral fluid using chicken eggs have been already developed (Nakanishi, et al. (eds.), 1993, "Shinkei-kagaku Kenkyu-no Sentan-gijutu Protocol III (High Technology Protocol III of Neuroscience Research), Molecular Neurocyte Physiology, Koseisha, Osaka, pp. 153-172). Specifically, for example, fertilized eggs are placed in an incubator and incubated for 9 to 12 days at 37 to 38°C to grow embryos. Sendai virus vector or RNP complex is inoculated together with vectors expressing envelope proteins into chorioallantoic cavity of eggs, and cultured for several days to proliferate the virus. Conditions such as culture duration may be varied depending on the type of recombinant Sendai virus used. Subsequently, chorioallantoic fluid comprising the virus is recovered. Separation and purification of Sendai virus vector can be performed according to the standard methods (Tashiro, M., "Virus Experiment Protocols", Nagai and Ishihama (eds.), Medicalview, pp. 68-73 (1995)).

[0041] As a vector to express envelope proteins, complexes of this invention or viral vectors themselves comprising complexes of this invention may be used. For example, when two types of RNP complexes in which the envelope gene deficient from the viral genome is different are transferred to the same cell, the envelope protein deficient in one RNP complex is supplied by the expression of the other complex to complement each other, thereby leading to the formation of infectious virus particles and activation of replication cycle to amplify the virus. That is, when two or more types of RNP complexes of this invention or viral vectors comprising these complexes are inoculated to cells in combinations so as to complement each other's envelope proteins, mixtures of viral vectors deficient in respective envelope proteins can be produced on a large scale and at a low cost. Mixed viruses thus produced are useful for the production of vaccines and such. Due to the deficiency of envelope genes, these viruses have a smaller genome size compared to the complete virus, so they can harbor a long foreign gene. Also, since these originally non-infectious viruses are extracellularly diluted, and its difficult to retain their coinfection, they become sterile, which is advantageous in managing their release to the environment.

[0042] Preparation of RNP of this invention from a virus can be carried out, for example, using the ultracentrifugation method as follows. Triton X-100 is added to a filtration fluid comprising virus particles to make the final concentration 0.5%, and the mixture is allowed to stand at room temperature for 10 to 15 min. The supernatant thus obtained is

layered on a 10 to 40% sucrose density gradient, and centrifuged at 20,000 to 30,000 rpm for 30 min to recover RNP-comprising fractions.

[0043] Alternatively, the virus is dissolved in a mixture containing 0.6% NP40, 1% sodium deoxycholate, 1 M KCl, 10 mM β -mercaptoethanol, 10 mM Tris-HCl (pH 7.4) and 5 mM EDTA (final concentrations), allowed to stand at 20°C for 20 min, and then centrifuged at 11,000 x g for 20 min. Supernatant comprising RNP is layered on 50% glycerol comprising 0.2% NP40, 30 mM NaCl, 10 mM Tris-HCl and 1 mM EDTA, and centrifuged at 39,000 rpm for 2 h at 4°C to recover precipitates. RNP complex contained in the precipitates can be purified by dispersing the precipitates again in a solution containing 0.5% Triton X-100, layering the dispersion on a 10 to 40% sucrose density gradient, and centrifuging it at 20,000 to 30,000 rpm for 30 min to recover a single band containing a highly purified RNP.

[0044] Complexes of this invention can be appropriately diluted, for example, with physiological saline and phosphate-buffered physiological saline (PBS) to prepare a composition. When complexes of this invention are proliferated in chicken eggs and such, the composition can include chorioallantoic fluid. Compositions comprising complexes of this invention may contain physiologically acceptable media such as deionized water, 5% dextrose aqueous solution, and so on, and, furthermore, other stabilizers and antibiotics may also be contained.

[0045] Once RNP-comprising RNA inserted with a foreign gene is prepared, it can be transferred to target cells using gene transfer reagents. As gene transfer reagents, cationic lipids or cationic polymers are preferred.

[0046] Cationic lipids include compounds represented by Formula (I) in Published Japanese Translation of International Publication No. Hei 5-508626. Preferably, cationic lipids are synthetic lipidic compounds. Cationic lipids may be also diether or diester compounds, preferably aliphatic ethers. Specific examples are the following compounds:

DOGS (Transfectam™) or DOTMA (Lipofectin™) (diether compound),
 DOTAP (diester compound),
 DOPE (dioleoylphosphatidylethanolamine),
 DOPC (dioleoylphosphatidylcholine),
 DPRI Rosenthal inhibitor (RI) (dipalmitoyl derivative of DL-2,3-distearoyloxypropyl (dimethyl) β -hydroxyethylammonium bromide (Sigma), and
 DORI (dioleoyl derivative of the above compound).

[0047] Cationic polymers are cationic high molecular compounds, preferably synthetic molecules. Specific examples are polylysine, aliphatic polyamines, polyethyleneimine, etc.

[0048] Complexes of this invention can be mixed with the above-described cationic lipids or cationic polymers to prepare compositions for gene transfer. This composition for gene transfer can be appropriately combined with a medium such as physiological saline, and solutes such as salts, stabilizers, etc. By adding the composition for gene transfer of this invention to cells, the complex of this invention can be transferred into the cells to express the gene from RNA contained in the complex.

[0049] Gene therapy is enabled when a therapeutic gene is used as the foreign gene. In the application of complexes of this invention to gene therapy, it is possible to express a foreign gene with which treatment effects are expected or an endogenous gene the supply of which is insufficient in the patient's body, by either direct or indirect (ex vivo) administration of the complex. There is no particular limitation on the type of foreign gene, and in addition to nucleic acids encoding proteins, they may be nucleic acids encoding no proteins, such as an antisense or ribozyme. In addition, when genes encoding antigens of bacteria or viruses involved in infectious diseases are used as foreign genes, immunity can be induced in animals by administering these genes to the animals. That is, these genes can be used as vaccines.

[0050] When using as vaccines, they may be applicable for, for example, cancers, infectious diseases and other general disorders. For example, as a cancer treatment, it is possible to express genes with therapeutic effects on tumor cells or antigen presenting cells (APC) such as DC cells. Examples of such genes are those encoding the tumor antigen Muc-1 or Muc-1 like mutin tandem repeat peptide (US Patent No. 5,744,144), melanoma gp100 antigen, etc. Such treatments with genes have been widely applied to cancers in the mammary gland, colon, pancreas, prostate, lung, etc. Combination with Gytokines to enhance adjuvant effects is also effective in gene therapy. Examples of such genes are i) single-chain IL-12 in combination with IL-2 (Proc. Natl. Acad. Sci. USA 96 (15): 8591-8596, ii) interferon- γ in combination with IL-2 (US Patent No. 5,798,100), iii) granulocyte colony-stimulating factor (GM-CSF) used alone, and iv) GM-CSF aiming at the treatment of brain tumor in combination with IL-4 (J. Neurosurgery, 90 (6), 1115-1124 (1999)), etc.

[0051] Examples of genes used for the treatment of infectious diseases are those encoding the envelope protein of the virulent strain H5NI type of influenza virus, the envelope chimera protein of Japanese encephalitis virus (Vaccine, vol. 17, No. 15-16, 1869-1882 (1999)), the HIV gag or SIV gag protein of AIDS virus (J. Immunology (2000), vol. 164, 4968-4978), the HIV envelope protein, which is incorporated as a oral vaccine encapsulated in polylactate- glycol copolymer microparticles for administration (Kaneko, H. et al., Virology 267, 8-16 (2000)), the B subunit (CTB) of

cholera toxin (Arakawa, T. et al., Nature Biotechnology (1998) 16 (10): 934-8; Arakawa, T. et al., Nature Biotechnology (1998) 16 (3): 292-297), the glycoprotein of rabies virus (Lodmell, D. L. et al., 1998, Nature Medicine 4 (8): 949-52), and the capsid protein L1 of human papilloma virus 6 causing cervical cancer (J. Med. Virol., 60, 200-204 (2000)).

[0052] Gene therapy may also be applied to general disorders. For example, in the case of diabetes, the expression of insulin peptide fragment by inoculation of plasmid DNA encoding the peptide has been performed in type I diabetes model animals (Coon, B. et al., J. Clin. Invest., 1999, 104 (2): 189-94).

Brief Description of the Drawings

[0053] Figure 1 is a photograph showing an analytical result of the expression of F protein via a Cre-loxP-inducible expression system by Western blotting. It shows the result of detecting proteins on a transfer membrane cross-reacting to the anti-SeV-F antibody by chemiluminescence method.

[0054] Figure 2 indicates a diagram showing an analytical result of cell-surface display of F protein the expression of which was induced by the Cre-loxP system. It shows results of flow cytometry analysis for LLC-MK2/F7 with the anti-SeV-F antibody.

[0055] Figure 3 indicates a photograph showing the result confirming cleavage of the expressed F protein by trypsin using Western blotting.

[0056] Figure 4 indicates photographs showing the result confirming cell-surface expression of HN in an experiment of cell-surface adsorption onto erythrocytes.

[0057] Figure 5 indicates photographs showing the result obtained by an attempt to harvest the deficient viruses by using cells expressing the deficient protein. It was revealed that the expression of F protein by the helper cell line was stopped rapidly by the vaccinia viruses used in the reconstitution of F-deficient SeV.

1. LLC-MK2 and CV-1 represent cell lysates from the respective cell types alone.

2. LLC-MK2/F+ad and CV-1/F+ad represent cell lysates from the respective cells that have been subjected to the induction of expression and to which adenovirus AxCANCre has been added.

3. LLC-MK2/F-ad and CV-1/F-ad represent cell lysates from the respective cell lines in which the F gene but no adenovirus AxCANCre has been introduced.

4. LLC-MK2/F+ad 3rd represents a cell lysate from cells in which the expression was induced by adenovirus AxCANCre and which were then further passaged 3 times.

5. 1d and 3d respectively indicate one day and three days after the induction of expression.

6. Vac1d and Vac3d respectively indicate cells one day and three days after the infection of vaccinia virus.

7. Ara1d and Ara3d respectively indicate cells one day and three days after the addition of AraC.

8. CHX 1d and CHX 3d respectively indicate cells one day and three days after the addition of protein synthesis inhibitor cycloheximide.

[0058] Figure 6 indicates photographs showing the result that was obtained by observing GFP expression after GFP-comprising F-deficient SeV cDNA (pSeV18*/ΔF-GFP) was transfected into LLC-MK2 cells in which F was not expressed (detection of RNP). In a control group, the F gene was shuffled with the NP gene at the 3' end, and then, SeV cDNA (F-shuffled SeV), in which GFP had been introduced into the F-deficient site, was used. The mark "all" indicates cells transfected with plasmids directing the expression of the NP gene, P gene, and L gene (pGEM/NP, pGEM/P, and pGEM/L) together with SeV cDNA at the same time; "cDNA" indicates cells transfected with cDNA (pSeV18*/ΔF GFP) alone. For RNP transfection, P0 cells expressing GFP were collected; the cells (10^7 cells/ml) were suspended in OptiMEM (GIBCO BRL); 100 μl of lysate prepared after treating three times with freeze-thaw cycles was mixed with 25 μl of cationic liposome DOSPER (Bdehringer Mannheim) and allowed to stand still at room temperature for 15 minutes; and the mixture was added to cells (+ad) in which the expression of F had been induced to achieve the RNP transfection. Cells expressing Cre DNA recombinase, in which no recombinant adenovirus had been introduced, were used as a control group of cells (-ad). The result showed that GFP was expressed depending on the RNP formation of SeV in P0 in LLC-MK2 cells; and the F-deficient virus was amplified depending on the induction of expression of F in P1.

[0059] Figure 7 indicates photographs showing the result that was obtained by studying whether functional RNP reconstituted with F-deficient genomic cDNA could be rescued by the F-expressing helper cells and form the infective virion of the deficient virus. RNP/o represents cells overlaid with RNP; RNP/t represents cells that was transfected with RNP.

[0060] Figure 8 indicates photographs showing the evidence for the F-expressing cell-specific growth of the F-deficient virus. The lysate comprising functional RNP constructed from the genome lacking the gene was lipofected to the F-expressing cells as described in Example 2; and the culture supernatant was then recovered. This culture supernatant was added to the medium of the F-expressing cells to achieve the infection; on the third day, the culture supernatant was recovered and concurrently added to both F-expressing cells and cells that had not expressed F; and then the

cells were cultured in the presence or absence of trypsin for three days. The result is shown here. The viruses were amplified only in the presence of trypsin in the F-expressing cells.

[0061] Figure 9 indicates photographs showing evidence for specific release of the F-deficient viruses to the culture supernatant after the introduction into F-expressing cells. The lysate comprising functional RNP constructed from the genome lacking the gene was lipofected to the F-expressing cells as described in Example 2 and then the culture supernatant was recovered. This culture supernatant was added to the medium of the F-expressing cells to achieve the infection; on the third day, the culture supernatant was recovered and concurrently added to both F-expressing cells and cells that did not express F; and then the cells were cultured in the presence or absence of trypsin for three days. The bottom panel shows the result with supernatant of the cells that did not express F.

[0062] Figure 10 indicates photographs showing the result obtained by recovering viruses from the culture supernatant of the F-expressing cells, extracting the total RNA and performing Northern blot analysis using F and HN as probes to verify the genomic structure of virion recovered from the F-deficient cDNA. In the viruses recovered from the F-expressing cells, the HN gene was detected but the F gene was not detectable; and thus it was clarified that the F gene was not present in the viral genome.

[0063] Figure 11 indicates photographs showing the result of RT-PCR, which demonstrates that the GFP gene is present in the locus where F had been deleted, as in the construct of the cDNA. 1: +18-NP, for the confirmation of the presence of +18 NotI site. 2: M-GFP, for the confirmation of the presence of the GFP gene in the F gene-deficient region. 3: F gene, for the confirmation of the presence of the F gene. The genomic structures of wild type SeV and F-deficient GFP-expressing SeV are shown in the top panel. It was verified that the GFP gene was present in the F-deficient locus, +18-derived NotI site was present at the 3' end of NP and the F gene was absent in any part of the RNA genome.

[0064] Figure 12 indicates photographs that were obtained by the immuno-electron microscopic examination with gold colloid-bound IgG (anti-F, anti-HN) specifically reacting to F or HN of the virus. It was clarified that the spike-like structure of the virus envelope comprised F and HN proteins.

[0065] Figure 13 indicates diagrams showing the result of RT-PCR, which demonstrates that the structures of genes except the GFP gene were the same as those from the wild type.

[0066] Figure 14 indicates photographs showing the result obtained by examining the F-deficient virus particle morphology by electron microscopy. Like the wild-type virus particles, the F-deficient virus particles had helical RNP structure and spike-like structure inside.

[0067] Figure 15 indicates photographs showing the result of *in vitro* gene transfer to a variety of cells using an F-deficient SeV vector with a high efficiency.

[0068] Figure 16 indicates diagrams showing the analytical result obtained after the introduction of the F-deficient SeV vector into primary bone marrow cells from mouse (BM c-kit⁺/-). Open bars represent PE-positive/GFP-negative; closed bars represent PE-positive/GFP-positive.

[0069] Figure 17 indicates photographs showing the result of *in vivo* administration of the vector into the rat cerebral ventricle.

[0070] Figure 18 indicates photographs showing the result obtained by using the culture supernatant comprising F-deficient SeV viruses recovered from the F-expressing cells to infect LLC-MK2 cells that do not express F, culturing the cells in the presence or absence of trypsin for three days to confirm the presence of viruses in the supernatant by HA assay.

[0071] Figure 19 is a photograph showing the result obtained by conducting HA assay of chorioallantoic fluids after a 2-day incubation of embryonated chicken egg that had been inoculated with chorioallantoic fluid (lanes 11 and 12) from HA-positive embryonated eggs in Figure 18B.

[0072] Figure 20 indicates photographs showing the result obtained by examining the virus liquid, which is HA-positive and has no infectivity, by immuno-electron microscopy. The presence of the virus particles was verified and it was found that the virion envelope was reactive to antibody recognizing HN- protein labeled with gold colloid, but not reactive to antibody recognizing F protein labeled with gold colloid.

[0073] Figure 21 indicates photographs showing the result of transfection of F-deficient virus particles into cells.

[0074] Figure 22 indicates photographs showing the result of creation of cells co-expressing F and HN, which were evaluated by Western blotting. LLC/VacT7/pGEM/FHN represents cells obtained by transfecting vaccinia-infected LLC-MK2 cells with pGEM/FHN plasmid; LLC/VacT7 represents vaccinia-infected LLC-MK2 cells. LLC/MK2/FHNmix represents LLC-MK2 cells in which the F and HN genes were introduced but not cloned. LLC/FHN represents LLC-MK2 cells in which the F and HN genes were introduced and the expression was induced by adenovirus (after 3 days); 1-13, 2-6, 2-16, 3-3, 3-18, 3-22, 4-3 and 5-9 are cell-line numbers (names) in the cloning.

[0075] Figure 23 indicates photographs showing the result for the confirmation of virus generation depending on the presence or absence pGEM/FHN. FHN-deficient GFP-expressing SeV cDNA, pGBM/NP, pGEM/P, pGEM/L, and pGEM/FHN were mixed and introduced into LLC-MK2 cells. 3 hours after the gene transfer, the medium was changed with MEM containing AraC and trypsin and then the cells were further cultured for three days. 2 days after the gene

transfer, observation was carried out with a stereoscopic fluorescence microscope to evaluate the difference depending on the presence or absence of pGEM/FHN, and the virus generation was verified based on the spread of GFP-expressing cells. The result is shown here. When pGEM/FHN was added at the time of reconstitution, the spread of GFP-expressing cells was recognized; but when no pGEM/FHN was added, the GFP expression was observable merely in a single cell.

[0076] Figure 24 indicates photographs showing the result of reconstitution by RNP transfection and growth of FHN-deficient viruses. On the third day after the induction of expression, cells co-expressing FHN (12 wells) were lipofected by using P0 RNP overlay or DOSPER, and then GFP was observed after 4 days. When RNP transfection was conducted, the harvest of viruses was successful for P1 FHN-expressing cells as was for the F-deficient ones (top). The growth of the FHN-deficient viruses was verified after inoculating a liquid comprising the viruses to cells in which the expression of FHN protein was induced 6 hours or more after the infection with Ade/Cre (bottom panel).

[0077] Figure 25 indicates photographs showing the result obtained after inoculating the liquid comprising viruses reconstituted from FHN-deficient GFP-expressing CDNA to LLC-MK2, LLC-MK2/F, LLC-MK2/HN, and LLC-MK2/FHN and culturing them in the presence or absence of the trypsin. The spread of cells expressing GFP protein was verified 3 days after the culture. The result is shown here. The expansion of GFP was observed only with LLC-MK2/FHN, and thus it was verified that the virus contained in the liquid was grown in a manner specific to FHN co-expression and dependent on trypsin.

[0078] Figure 26 is a photograph showing the result where the confirmation was carried out for the genomic structure of RNA derived from supernatant of the FHN-expressing cells.

[0079] Figure 27 is a photograph showing the result where the confirmation was carried out for the genomic structure of RNA derived from supernatant of the F-expressing cells infected with the FHN-deficient viruses.

[0080] Figure 28 is a diagram showing inactivation of vaccinia virus and T7 activity when psoralen concentration was varied in psoralen/UV irradiation.

[0081] Figure 29 is a diagram showing inactivation of vaccinia virus and T7 RNA polymerase activity when the duration of UV irradiation was varied in psoralen/UV irradiation.

[0082] Figure 30 indicates photographs showing a cytotoxicity (CPE) of vaccinia virus after psoralen/UV irradiation. 3×10^5 LLC-MK2 cells were plated on a 6-well plate. After culturing overnight, the cells were infected with vaccinia virus at $\text{moi}=2$. After 24 hours, CPE was determined. The result of CPE with mock-treatment of vaccinia virus is shown in A; CPE after the treatment with vaccinia virus for 15, 20, or 30 minutes are shown in B, C, and D, respectively.

[0083] Figure 31 is a diagram indicating the influence of duration of UV treatment of vaccinia virus on the reconstitution efficiency of Sendai virus.

[0084] Figure 32 is a diagram indicating the titer of vaccinia virus capable of replicating that remained in the cells after the reconstitution experiment of Sendai virus.

[0085] Figure 33 is a photograph showing a result of Western blot analysis using anti-VSV-G antibody.

[0086] Figure 34 indicates a diagram showing results of flow cytometry analysis using anti-VSV-G antibody. It shows the result of analysis of LLC-MK2 cell line (L1) for the induction of VSV-G expression on the fourth day after AxCANCre infection ($\text{moi}=0, 2.5, 5$). Primary antibody used was anti-VSV-G antibody (MoAb I-1); secondary antibody was FITC-labeled anti-mouse Ig.

[0087] Figure 35 indicates photographs showing a result where supernatants were recovered after the infection with altered amounts of AxCANCre ($\text{MOI}=0, 1.25, 2.5, 5, 10$) and a constant amount of pseudo-type Sendai virus having a F gene-deficient genome, and further the supernatants were used to infect cells before VSV-G induction (-) and after induction (+), and cells expressing GFP were observed after 5 days.

[0088] Figure 36 indicates photographs showing the result obtained for the time course of virus production amount.

[0089] Figure 37 indicates photographs showing the result obtained by examining whether the infectivity is influenced by the treatment of pseudo-type Sendai virus having the F gene-deficient genome, which was established with the VSV-G-expressing cell line, and FHN-deficient Sendai virus treated with anti-VSV antibody.

[0090] Figure 38 indicates photographs showing the result where the expression of the GFP gene was tested as an index to determine the presence of production of the pseudo-type virus having VSV-G in its capsid after the infection of VSV-G gene-expressing cells LLCG-L1 with F and HN-deficient Sendai virus comprising the GFP gene.

[0091] Figure 39 indicates photographs showing the result confirming that viruses grown in the VSV-G gene-expressing cells were deficient in F and HN genes by Western analysis of protein in the extract of infected cells.

[0092] Figure 40 indicates photographs showing the result for the observation of GFP-expressing cells under a fluorescence microscope.

[0093] Figure 41 is a diagram showing the improvement in efficiency for the reconstitution of SeV/ Δ F-GFP by the combined use of the envelope-expressing plasmid and cell overlay. Considerable improvement was recognized at d3 to d4 (day 3 to day 4) of P0 (prior to passaging).

[0094] Figure 42 is a diagram showing the result where treatment conditions were evaluated for the reconstitution of SeV/ Δ F-GFP by the combined use of the envelope-expressing plasmid and cell overlay. GFP-positive cells repre-

sent the amount of virus reconstituted.

[0095] Figure 43 is a diagram showing the result where the rescue of F-deficient Sendai viruses from cDNA was tested. It shows the improvement in efficiency for the reconstitution of SeV/ΔF-GFP by the combined use of the envelope-expressing plasmid and cell overlay. All the tests were positive on the seventh day. However, the efficiency was evaluated on the third day where the probability of success was midrange.

[0096] Figure 44 indicates photographs showing the result of lacZ expression by LacZ-comprising F-deficient Sendai virus vector comprising no GFP.

[0097] Figure 45 indicates diagrams showing subcloning of Sendai virus genomic cDNA fragment (A) and structures of 5 Sendai virus genomic cDNAs constructed with newly introduced NotI site (B).

[0098] Figure 46 is a diagram showing structures of plasmids to be used for cloning to add NotI site, transcription initiation signal, intervening sequence, and transcription termination signal into SEAP.

[0099] Figure 47 indicates photographs showing the result of plaque assay of each Sendai virus vector. It shows partial fluorescence image in the plaque assay obtained by LAS1000.

[0100] Figure 48 is a diagram showing the result where altered expression levels of reporter gene (SEAP) were compared with one another among the respective Sendai virus vectors. The data of SeV18+/SEAP was taken as 100 and the respective values were indicated relative to it. It was found that the activity, namely the expression level, was decreased as the SEAP gene was placed more downstream.

[0101] Figure 49 indicates microscopic photographs showing the expression of GFP in P1 cells co-expressing FHN.

[0102] Figure 50 indicates photographs showing the result of Western blot analysis of the extracts from cells infected with VSV-G pseudo-type SeV/ΔF:GFP using anti-F antibody (anti-F), anti-HN antibody (anti-HN), and anti-Sendai virus antibody (anti-SeV).

[0103] Figure 51 indicates photographs showing GFP fluorescence from F- and HN-deficient cells infected with VSV-G pseudo-type SeV in the presence or absence of a neutralizing antibody (VGV antibody).

[0104] Figure 52 indicates photographs showing results of Western analysis for VSV-G pseudo-type Sendai viruses having F gene-deficient or F gene- and HN gene-deficient genome, which were fractionated by density gradient ultracentrifugation.

[0105] Figure 53 indicates photographs showing hemagglutination test mediated with Sendai viruses having F gene-deficient genome, or VSV-G pseudo-type Sendai viruses having F gene-deficient or F gene- and HN gene-deficient genome.

[0106] Figure 54 indicates diagrams showing the specificity of infection to culture cells of Sendai virus having F gene-deficient genome or VSV-G pseudo-type Sendai virus.

[0107] Figure 55 indicates photographs showing the confirmation of the structures of NGF-expressing F-deficient Sendai virus (NGF/SeV/ΔF).

[0108] Figure 56 is a diagram showing the activity of NGF expressed by the NGF-comprising cells infected with F-deficient SeV. With the initiation of culture, diluted supernatant of SeV-infected cells or NGF protein (control) was added to a dissociated culture of primary chicken dorsal root ganglion (DRG) neurons. After three days, the viable cells were counted by using mitochondrial reduction activity as an index (n=3). The quantity of culture supernatant added corresponded to 1000-fold dilution.

[0109] Figure 57 indicates photographs showing the activity of NGF expressed by the NGF-comprising cells infected with F-deficient SeV. With the initiation of culture, diluted supernatant of SeV-infected cells or NGF protein (control) was added to a dissociated culture of primary chicken dorsal root ganglion (DRG) neurons. After three days, the samples were observed under a microscope,

A) control (without NGF);

B) addition of NGF protein (10 ng/mL);

C) addition of culture supernatant (100-fold diluted) of NGF/SeV infected cells;

D) addition of culture supernatant (100-fold diluted) of NGF/SeV infected cells;

E) addition of culture supernatant (100-fold diluted) of NGF/SeV/ΔF infected cells, and;

F) addition of culture supernatant (100-fold diluted) of NGF/SeV/ΔF-GFP infected cells.

[0110] Figure 58 is a photograph showing moi of Ad-Cre and the expression level of F protein.

[0111] Figure 59 indicates photographs showing the expression of LLC-MK2/F by Adeno-Cre.

[0112] Figure 60 is a photograph showing the durability of expression over the passages.

[0113] Figure 61 indicates photographs showing the localization of F protein over the passages.

[0114] Figure 62 is a diagram showing the correlation between GFP-CIU and anti-SeV-CIU.

Best Mode for Carrying out the Invention

[0115] The present invention is illustrated in detail below with reference to Examples, but is not to be construed as being limited thereto.

[Example 1] Construction of F-deficient Sendai virus

<1> Construction of F-deficient SeV genomic cDNA and F-expressing plasmid

[0116] The full-length genomic cDNA of Sendai virus (SeV), pSeV18⁺b(+) (Hasan, M. K. et al., 1997, J. General Virology 78: 2813-2820). ("pSeV18⁺ b(+)") is also referred to as "pSeV18⁺") was digested with SphI/KpnI, and the resulting fragment (14673 bp) was recovered and cloned into pUC18, which was named plasmid pUC18/KS. The F-disrupted site was constructed on this pUC18/KS. The F gene disruption was performed by the combined use of PCR-ligation method, and as a result, the ORF for the F gene (ATG-TGA=1698 bp) was removed; thus atgcatgccggcagatga (SEQ ID NO: 1) was ligated to it to construct the F-deficient SeV genomic cDNA (pSeV18⁺/ΔF). In PCR, a PCR product generated by using a primer pair (forward: 5'-gttgagtactgcaagagc/SEQ ID NO: 2, reverse: 5'-ttgcccgcgatgcatgttcccaaggagagatgttgcaacc/SEQ ID NO: 3) was ligated upstream of F and another PCR product generated by using a primer pair (forward: 5'-atgcatgccggcagatga/SEQ ID NO: 4, reverse: 5'-tgggtgaatgagagaatcagc/SEQ ID NO: 5) was ligated downstream of the F gene at EcoT22I site. The resulting plasmid was digested with SacI and Sall, and then the fragment (4931 bp) spanning the region comprising the site where F is disrupted was recovered and cloned into pUC18 to generate pUC18/dFSS. This pUC18/dFSS was digested with DraIII. The resulting fragment was recovered and substituted with a DraIII fragment from the region comprising the F gene of pSeV18⁺; and the ligation was carried out to generate plasmid pSeV18⁺/ΔF.

[0117] Further, in order to construct a cDNA (pSeV18⁺/ΔF-GFP) in which the EGFP gene has been introduced at the site where F was disrupted, the EGFP gene was amplified by PCR. To set the EGFP gene with a multiple of 6 (Hausmann, S. et al., RNA 2, 1033-1045 (1996)), PCR was carried out with an NsiI-tailed primer (5'-atgcatatggtgatgcgggtttggcagtac: SEQ ID NO: 6) for the 5' end and an NgoMIV-tailed primer (5'-Tgccggctattattactgttacagctcgtc: SEQ ID NO: 7) for the 3' end. The PCR products were digested with restriction enzymes NsiI and NgoMIV, and then the fragment was recovered from the gel; the fragment was ligated at the site of pUC18/dFSS between NsiI and NgoMIV restriction enzyme sites where the disrupted F is located and the sequence was determined. A DraIII fragment comprising the EGFP gene was removed and recovered from the site, and substituted for a DraIII fragment in the region comprising the F gene of pSeV18⁺; then ligation was carried out to obtain plasmid pSeV18⁺/ΔF-GFP.

[0118] On the other hand, Cre-loxP-inducible expression plasmid for F gene expression was constructed by amplifying the SeV F gene by PCR, confirming the sequence, and inserting into the unique site SwaI of plasmid pCALNdlw (Arai et al., J. Virology 72, 1998, p1115-1121), in which the expression of gene products has been designed to be induced by Cre DNA recombinase, to obtain plasmid pCALNdlw/F.

<2> Preparation of helper cells inducing the expression of SeV-F protein

[0119] To recover infectious virus particles from F-deficient genome, a helper cell strain expressing SeV F protein was established. The cell utilized was LLC-MK2 cell that is commonly used for the growth of SeV and is a cell strain derived from monkey kidney. The LLC-MK2 cells were cultured in MEM containing 10% heat-treated inactivated fetal bovine serum (FBS), sodium penicillin G (50 units/ml), and streptomycin (50 μg/ml) at 37°C under 5% CO₂ gas. Because SeV-F gene product is cytotoxic, the above-mentioned plasmid pCALNdlw/F designed to induce the expression of F gene product through Cre DNA recombinase was introduced into LLC-MK2 cells by calcium phosphate method (mammalian transfection kit (Stratagene)) according to the gene transfer protocol.

[0120] 10 μg of plasmid pCALNdlw/F was introduced into LLC-MK2 cells grown to be 40% confluent in a 10-cm plate, and the cells were cultured in 10 ml of MEM containing 10% FBS at 37°C under 5% CO₂ for 24 hours in an incubator. After 24 hours, the cells were scraped off, and suspended in 10 ml medium; then the cells were plated on 5 dishes with 10-cm diameter (one plate with 5 ml; 2 plates with 2 ml; 2 plates with 0.2 ml) in MEM containing 10 ml of 10% FBS and 1200 μg/ml G418 (GIBCO-BRL) for the cultivation. The culture was continued for 14 days while the medium was changed at 2-day intervals, to select cell lines in which the gene has been introduced stably. 30 cell strains were recovered as G418-resistant cells grown in the medium by using cloning rings. Each clone was cultured to be confluent in 10-cm plates.

[0121] After the infection of each clone with recombinant adenovirus AXCANCre expressing Cre DNA recombinase, the cells were tested for the expression of SeV-F protein by Western blotting using anti-SeV-F protein monoclonal IgG (f236; J. Biochem. 123: 1064-1072) as follows.

[0122] After grown to be confluent in a 6-cm dish, each clone was infected with adenovirus AxCANCre at moi=3 according to the method of Saito et al., (Saito et al., Nucl. Acids Res. 23: 3816-3821 (1995); Arai, T. et al., J Virol 72, 1115-1121 (1998)). After the infection, the cells were cultured for 3 days. The culture supernatant was discarded and the cells were washed twice with PBS buffer, scraped off with a scraper and were collected by centrifugation at 1500x g for five minutes.

[0123] The cells are kept at -80°C and can be thawed when used. The cells collected were suspended in 150 µl PBS buffer, and equal amount of 2x Tris-SDS-BME sample loading buffer (0.625 M Tris, pH 6.8, 5% SDS, 25% 2-ME, 50% glycerol, 0.025% BPB; Owl) was added thereto. The mixture was heat-treated at 98°C for 3 minutes and then used as a sample for electrophoresis. The sample (1x 10⁵ cells/lane) was fractionated by electrophoresis in an SDS-polyacrylamide gel (Multi Gel 10/20, Daiichi Pure Chemicals). The fractionated proteins were transferred onto a PVDF transfer membrane (Immobilon-P transfer membranes; Millipore) by semi-dry blotting. The transfer was carried out under a constant current of 1 mA/cm² for 1 hour onto the transfer membrane that had been soaked in 100% methanol for 30 seconds and then in water for 30 minutes.

[0124] The transfer membrane was shaken in a blocking solution containing 0.05% Tween20 and 1% BSA (BlockAce; Snow Brand Milk Products) for one hour, and then it was incubated at room temperature for 2 hours with an anti-SeV-F antibody (f236) which had been diluted 1000-folds with a blocking solution containing 0.05% Tween 20 and 1% BSA. The transfer membrane was washed 3 times in 20 ml of PBS-0.1% Tween20 while being shaken for 5 minutes and then it was washed in PBS buffer while being shaken for 5 minutes. The transfer membrane was incubated at room temperature for one hour in 10 ml of peroxidase-conjugated anti-mouse IgG antibody (Goat anti-mouse IgG; Zymed) diluted 2000-fold with the blocking solution containing 0.05% Tween 20 and 1% BSA. The transfer membrane was washed 3 times with 20 ml of PBS-0.1% Tween20 while being shaken for 5 minutes, and then it was washed in PBS buffer while being shaken for 5 minutes.

[0125] Detections were carried out for proteins cross-reacting to the anti-SeV-F antibody on the transfer membrane by chemiluminescence method (ECL western blotting detection reagents; Amersham). The result is shown in Figure 1. The SeV-F expression specific to AxCANCre infection was detected to confirm the generation of LLC-MK2 cells that induce expression of a SeV-F gene product.

[0126] One of the several resulting cell lines, LLC-MK2/F7 cell, was analyzed by flow cytometry with an anti-SeV-F antibody (Figure 2). Specifically, 1x 10⁵ cells were precipitated by centrifugation at 15,000 rpm at 4°C for 5 minutes, washed with 200 µl PBS, and allowed to react in PBS for FACS (NIKKEN CHEMICALS) containing 100-fold diluted anti-F monoclonal antibody (f236), 0.05% sodium azide, 2% FCS at 4°C for 1 hour in a dark place. The cells were again precipitated at 15,000 rpm at 4°C for 5 minutes, washed with 200 µl PBS, and then allowed to react to FITC-labeled anti-mouse IgG (CAPPEL) of 1 µg/ml on ice for 30 minutes. Then the cells were again washed with 200 µl PBS, and then precipitated by centrifugation at 15,000 rpm at 4°C for 5 minutes. The cells were suspended in 1 ml of PBS for FACS and then analyzed by using EPICS ELITE (Coulter) argon laser at an excitation wavelength of 488 nm and at a fluorescence wavelength of 525 nm. The result showed that LLC-MK2/F7 exhibited a high reactivity to the antibody in a manner specific to the induction of SeV-F gene expression, and thus it was verified that SeV-F protein was expressed on the cell surface.

[Example 2] Confirmation of function of SeV-F protein expressed by helper cells

[0127] It was tested whether or not SeV-F protein, of which expression was induced by helper cells, retained the original protein function.

[0128] After plating on a 6-cm dish and grown to be confluent, LLC-MK2/F7 cells were infected with adenovirus AxCANCre at moi-3 according to the method of Saito et al. (described above). Then, the cells were cultured in MEM (serum free) containing trypsin (7.5 µg/ml; GIBCOBRL) at 37°C under 5% CO₂ in an incubator for three days.

[0129] The culture supernatant was discarded and the cells were washed twice with PBS buffer, scraped off with a scraper, and collected by centrifugation at 1500 x g for five minutes. The cleavage of expressed F protein by trypsin was verified by Western blotting as described above (Figure 3). SeV-F protein is synthesized as F0 that is a non-active protein precursor, and then the precursor is activated after being digested into two subunits F1 and F2 by proteolysis with trypsin. LLC-MK2/F7 cells after the induction of F protein expression thus, like ordinary cells, continues to express F protein, even after being passaged, and no cytotoxicity mediated by the expressed F protein was observed as well as no cell fusion of F protein-expressing cells was observed. However, when SeV-HN expression plasmid (pCAG/SeV-HN) was transfected into the F-expressing cells and the cells were cultured in MEM containing trypsin for 3 days, cell fusion was frequently observed. The expression of HN on the cell surface was confirmed in an experiment using erythrocyte adsorption onto the cell surface (Hematoadsorption assay; Had assay) (Figure 4). Specifically, 1% chicken erythrocytes were added to the culture cells at a concentration of 1 ml/dish and the mixture was allowed to stand still at 4°C for 10 minutes. The cells were washed 3 times with PBS buffer, and then colonies of erythrocytes on the cell surface were observed. Cell fusion was recognized for cells on which erythrocytes aggregated; cell fusion was found

to be induced through the interaction of F protein with HN; and thus it was demonstrated that F protein, the expression of which was sustained in LLC-MK2/F7, retained the original function thereof.

[Example 3] Functional RNP having F-deficient genome and formation of virions

[0130] To recover virions from the deficient viruses, it is necessary to use cells expressing the deficient protein. Thus, the recovery of the deficient viruses was attempted with cells expressing the deficient protein, but it was revealed that the expression of F protein by the helper cell line stopped rapidly due to the vaccinia viruses used in the reconstitution of F-deficient SeV (Figure 5) and thus the virus reconstitution based on the direct supply of F protein from the helper cell line failed. It has been reported that replication capability of vaccinia virus is inactivated, but the activity of T7 expression is not impaired by the treatment of vaccinia virus with ultraviolet light of long wavelengths (long-wave UV) in the presence of added psoralen (PLWUV treatment) (Tsung et al., J Virol 70, 165-171, 1996). Thus, virus reconstitution was attempted by using PLWUV-treated vaccinia virus (PLWUV-VacT7). UV Stratalinker 2400 (Catalog NO. 400676 (100V); Stratagene, La Jolla, CA, USA) equipped with five 15-Watt bulbs was used for ultraviolet light irradiation. The result showed that the expression of F protein was inhibited from the F-expressing cells used in the reconstitution, but vaccinia was hardly grown in the presence of araC after lysate from the cells reconstituted with this PLWUV-VacT7 was infected to the helper cells, and it was also found that the expression of F protein by the helper cell line was hardly influenced. Further, this reconstitution of wild type SeV using this PLWUV-VacT7 enables the recovery of viruses from even 10^3 cells, whereas by previous methods, this was not possible unless 10^5 or more cells were there, and thus the efficiency of virus reconstitution was greatly improved. Thus, reconstitution of F-deficient SeV virus was attempted by using this method.

<Reconstitution and amplification of F-deficient SeV virus>

[0131] The expression of GFP was observed after transfecting LLC-MK2 cells with the above-mentioned pSeV18⁺/ΔF-GFP in which the enhanced green fluorescent protein (EGFP) gene had been introduced as a reporter into the site where F had been disrupted according to the 6n rule in the manner as described below. It was also tested for the influence of the presence of virus-derived genes NP, P, and L that are three components required for the formation of RNP.

[0132] LLC-MK2 cells were plated on a 100-mm Petri-dish at a concentration of 5×10^6 cells/dish and were cultured for 24 hours. After the culture was completed, the cells were treated with psoralen and ultraviolet light of long wavelengths (365 nm) for 20 minutes, and the cells were infected with recombinant vaccinia virus expressing T7 RNA polymerase (Fuerst, T.R. et al., Proc. Natl. Acad. Sci. USA 83, 8122-8126 (1986)) at room temperature for one hour (moi=2) (moi=2 to 3; preferably moi=2). After the cells were washed 3 times, plasmids pSeV18⁺/ΔF-GFP, pGEM/NP, pGEM/P, and pGEM/L (Kato, A. et al., Genes cells 1, 569-579 (1996)) were respectively suspended in quantities of 12 μg, 4 μg, 2 μg, and 4 μg /dish in OptiMEM (GIBCO); SuperFect transfection reagent (1 μg DNA/5 μl SuperFect; QIAGEN) was added thereto; the mixtures were allowed to stand still at room temperature for 10 minutes; then they are added to 3 ml of OptiMEM containing 3% FBS; cells were added thereto and cultured. The same experiment was carried out using wild-type SeV genomic cDNA (pSeV(+)) (Kato, A. et al., Genes cells 1, 569-579 (1996)) as a control instead of pSeV18⁺/ΔF-GFP. After culturing for 3 hours, the cells were washed twice with MEM containing no serum, and then cultured in MEM containing cytosine β-D-arabinofuranoside (AraC, 40 μg/ml; Sigma) and trypsin (7.5 μg/ml; GIBCO) for 70 hours. These cells were harvested, and the pellet was suspended in OptiMEM (10^7 cells/ml). After freeze-and-thaw treatment was repeated 3 times, the cells were mixed with lipofection reagent DOSPER (Boehringer Mannheim) (10^6 cells/25 μl DOSPER) and allowed to stand still at room temperature for 15 minutes. Then F-expressing LLC-MK2/F7 cell line (10^6 cells /well in 12-well plate) was transfected, and the cells were cultured in MEM containing no serum (containing 40 μg/ml AraC and 7.5 μg/ml trypsin).

[0133] The result showed that the expression of GFP was recognized only when all the three components, NP, P, and L derived from the virus are present and the deficient virus RNP expressing foreign genes can be generated (Figure 6).

<Confirmation of F-deficient virions>

[0134] It was tested whether the functional RNP reconstituted by F-deficient genomic cDNA by the method as described above could be rescued by the F-expressing helper cells and form infective virions of F-deficient virus. Cell lysates were mixed with cationic liposome; the lysates were prepared by freeze/thaw from cells reconstituted under conditions in which functional RNP is formed (condition where pSeV18⁺/ΔF-GFP, pGEM/NP, pGEM/P, and pGEM/L are transfected at the same time) or conditions under which functional RNP is not formed (conditions in which two plasmids, pSeV18⁺/ΔF-GFP and pGEM/NP, are transfected) as described above; the lysates were lipofected into F-

expressing cells and non-expressing cells; the generation of virus particles was observed based on the expansion of the distribution of GFP-expressing cells. The result showed that the expansion of distribution of GFP-expressing cells was recognized only when the introduction to the F-expressing cells was carried out by using a lysate obtained under condition in which functional RNP is reconstituted (Figure 7). Furthermore, even in plaque assay, the plaque formation was seen only under the same conditions. From these results, it was revealed that functional RNPs generated from F-deficient virus genome were further converted into infective virus particles in the presence of F protein derived from F-expressing cells and the particles were released from the cells.

[0135] The demonstration of the presence of infective F-deficient virions in the culture supernatant was carried out by the following experiment. The lysate comprising the functional RNP constructed from the F gene deficient genome was lipofected to F-expressing cells as described in Example 2, and the culture supernatant was recovered. This culture supernatant was added to the medium of F-expressing cells to achieve the infection; on the third day, the culture supernatant was recovered and concurrently added to both F-expressing cells and cells that did not express F; and then the cells were cultured in the presence or absence of trypsin for three days. In F-expressing cells, viruses were amplified only in the presence of trypsin (Figure 8). It was also revealed that non-infectious virus particles were released into the supernatant of cells that do not express F (in the bottom panel of Figure 9) or from F-expressing cells cultured in the absence of trypsin. A summary of the descriptions above is as follows: the growth of F-deficient GFP-expressing viruses is specific to F-expressing cells and depends on the proteolysis with trypsin. The titer of infective F-deficient Sendai virus thus grown ranged from 0.5×10^7 to 1×10^7 CIU/ml.

[Example 4] Analysis of F-deficient GFP-expressing virus

[0136] In order to confirm the genomic structure of virions recovered from F-deficient cDNA, viruses were recovered from the culture supernatant of the F-expressing cells, the total RNA was extracted and then Northern blot analysis was conducted by using F and HN as probes. The result showed that the HN gene was detectable, but the F gene was not detectable in the viruses harvested from the F-expressing cells, and it was clarified that the F gene was not present in the viral genome (Figure 10). Further, by RT-PCR GFP, it was confirmed that the gene was present in the deleted locus for F as shown in the construction of the cDNA (Figure 11) and that the structures of other genes were the same as those from the wild type. Based on the findings above, it was shown that no rearrangement of the genome had occurred during the virus reconstitution. In addition, the morphology of recovered F-deficient virus particles was examined by electron microscopy. Like the wild type virus, F-deficient virus particles had the helical RNP structure and spike-like structure inside (Figure 14). Further, the viruses were examined by immuno-electron microscopy with gold colloid-conjugated IgG (anti-F, anti-HN) specifically reacting to F or HN. The result showed that the spike-like structure of the envelope of the virus comprised F and HN proteins (Figure 12), which demonstrated that F protein produced by the helper cells was efficiently incorporated into the virions. The result will be described below in detail.

<Extraction of total RNA, Northern blot analysis, and RT-PCR>

[0137] Total RNA was extracted from culture supernatant obtained 3 days after the infection of F-expressing cell LLC-MK2/F7 with the viruses by using QIAamp Viral RNA mini kit (QIAGEN) according to the protocol. The purified total RNA (5 µg) was separated by electrophoresis in a 1% denaturing agarose gel containing formaldehyde, and then transferred onto a Hybond-N+ membrane in a vacuum blotting device (Amersham-Pharmacia). The prepared membrane was fixed with 0.05 M NaOH, rinsed with 2-fold diluted SSC buffer (Nacal Tesque), and then was subjected to pre-hybridization in a hybridization solution (Boehringer Mannheim) for 30 minutes; a probe for the F or HN gene prepared by random prime DNA labeling (DIG DNA Labeling Kit; Boehringer Mannheim) using digoxigenin (DIG)-dUTP (alkaline sensitive) was added thereto and then hybridization was performed for 16 hours. Then, the membrane was washed, and allowed to react to alkaline phosphatase-conjugated anti-DIG antibody (anti-digoxigenin-AP); the analysis was carried out by using a DIG detection kit. The result showed that the HN gene was detectable but the F gene was not detectable in the viruses harvested from the F-expressing cells, and it was clarified that the F gene was not present in the viral genome (Figure 10).

[0138] Further, detailed analysis was carried out by RT-PCR. In the RT-PCR, first strand cDNA was synthesized from the purified virus RNA by using SUPERScript II Preamplification System (Gibco BRL) according to the protocol; the following PCR condition was employed with LA PCR kit (TAKARA ver2.1): 94°C/3 min; 30 cycles for the amplification of 94°C/45 sec, 55°C/45 sec, 72°C/90 sec; incubation at 72°C for 10 minutes; then the sample was electrophoresed in a 2% agarose gel at 100 v for 30 minutes, the gel was stained with ethidium bromide for a photographic image. Primers used to confirm the M gene and EGFP inserted into the F-deficient site were forward 1: 5'-atcagagacctgcgacaatgc (SEQ ID NO: 8) and reverse 1: 5'-aagtcgtgctgcttcctggtgg (SEQ ID NO: 9); primers used to confirm EGFP inserted into the F-deficient site and the HN gene were forward 2: 5'-acaaccactacctgagcaccagtc (SEQ ID NO: 10) and reverse 2: 5'-gcctaacacatccagagatcg (SEQ ID NO: 11); and the junction between the M gene and HN gene was confirmed by

using forward 3: 5'-acattcatgagtcagctcgc (SEQ ID NO: 12) and reverse 2 primer (SEQ ID NO: 11). The result showed that the GFP gene was present in the deficient locus for F as shown in the construction of the cDNA (Figure 11) and that the structures of other genes were the same as those from the wild type (Figure 13). From the findings shown above, it is clarified that no rearrangement of the genome had resulted during the virus reconstitution.

<Electron microscopic analysis with gold colloid-conjugated immunoglobulin>

[0139] The morphology of recovered F-deficient virus particles were examined by electron microscopy. First, culture supernatant of cells infected with the deficient viruses was centrifuged at 28,000 rpm for 30 minutes to obtain a virus pellet; then the pellet was re-suspended in 10-fold diluted PBS at a concentration of 1×10^9 HAU/ml; one drop of the suspension was dropped on a microgrid with a supporting filter and then the grid was dried at room temperature; the grid was treated with PBS containing 3.7% formalin for 15 minutes for fixation and then pre-treated with PBS solution containing 0.1% BSA for 30 minutes; further, anti-F monoclonal antibody (f236) or anti-HN monoclonal antibody (Miura, N. et al., Exp. Cell Res. (1982) 141: 409-420) diluted 200-folds with the same solution was dropped on the grid and allowed to react under a moist condition for 60 minutes. Subsequently, the grid was washed with PBS, and then gold colloid-conjugated anti-mouse IgG antibody diluted 200-folds was dropped and allowed to react under a moist condition for 60 minutes. Subsequently, the grid was washed with PBS and then with distilled sterile water, and air-dried at room temperature; 4% uranium acetate solution was placed on the grid for the staining for 2 minutes and the grid was dried; the sample was observed and photographed in a JEM-1200EXII electron microscope (JEOL.). The result showed that the spike-like structure of the envelope of the virus comprised F and HN proteins (Figure 12), which demonstrated that F protein produced by the helper cells was efficiently incorporated into the virions. In addition, like the wild type virus, F-deficient virus particles had a helical RNP structure and a spike-like structure inside (Figure 14).

[Example 5] High-efficiency gene transfer to a variety of cells via F-deficient SeV vector in vitro

<Introduction into primary culture cells of rat cerebral cortex nerve cells>

[0140] Primary culture cells of rat cerebral cortex neurons were prepared and cultured as follows: an SD rat (SPF/VAF Crj : CD, female, 332 g, up to 9-week old; Charles River) on the eighteenth day of pregnancy was deeply anesthetized by diethyl ether, and then euthanized by bloodletting from axillary arteries. The fetuses were removed from the uterus after abdominal section. The cranial skin and bones were cut and the brains were taken out. The cerebral hemispheres were transferred under a stereoscopic microscope to a working solution DMEM (containing 5% horse serum, 5% calf serum and 10% DMSO); they were sliced and an ice-cold papain solution (1.5 U, 0.2 mg of cysteine, 0.2 mg of bovine serum albumin, 5 mg glucose, DNase of 0.1 mg/ml) was added thereto; the solution containing the sliced tissues was incubated for 15 minutes while shaking by inverting the vial every 5 minutes at 32°C. After it was verified that the suspension became turbid enough and the tissue sections became translucent, the tissue sections were crushed into small pieces by pipetting. The suspension was centrifuged at 1200 rpm at 32°C for 5 minutes, and then the cells were re-suspended in B27-supplemented neural basal medium (GibcoBRL, Burlington, Ontario, Canada). The cells were plated on a plate coated with poly-d-lysine (Becton Dickinson Labware, Bedford, MA, U.S.A.) at a density of 1×10^5 cells/dish and then cultured at 37°C under 5% CO₂.

[0141] After the primary culture of nerve cells from cerebral cortex (5×10^5 /well) were cultured for 5 days, the cells were infected with F-deficient SeV vector (moi=5) and further cultured for three days. The cells were fixed in a fixing solution containing 1% paraformaldehyde, 5% goat serum, and 0.5% Triton-X at room temperature for five minutes. Blocking reaction was carried out for the cells by using BlockAce (Snow Brand Milk Products) at room temperature for 2 hours, and then incubated with 500-fold diluted goat anti-rat microtubule-associated protein 2 (MAP-2) (Boehringer) IgG at room temperature for one hour. Further, the cells were washed three times with PBS(-) every 15 minutes and then were incubated with cys3-conjugated anti-mouse IgG diluted 100-folds with 5% goat serum/PBS at room temperature for one hour. Further, after the cells were washed three times with PBS(-) every 15 minutes, Vectashield mounting medium (Vector Laboratories, Burlingame, U.S.A.) was added to the cells; the cells, which had been double-stained with MAP-2 immuno staining and GFP fluorescence, were fluorescently observed by using a confocal microscope (Nippon Bio-Rad MRC 1024, Japan) and an inverted microscope Nikon Diaphot 300 equipped with excitation band-pass filter of 470-500-nm or 510-550-nm. The result showed that GFP had been introduced in nearly 100% nerve cells that were MAP2-positive (Figure 15).

<Introduction into normal human cells>

[0142] Normal human smooth-muscle cells, normal human hepatic cells, and normal human pulmonary capillary endothelial cells (Cell Systems) were purchased from DAINIPPON PHARMACEUTICAL and were cultured with SFM

CS-C medium kit (Cell Systems) at 37°C under 5% CO₂ gas.

[0143] Human normal cells, such as normal human smooth-muscle cells (Figure 15, Muscle), normal human hepatic cells (Figure 15, Liver), and normal human pulmonary capillary endothelial cells (Figure 15, Lung), were infected with F-deficient SeV vector (m.o.i.=5), and then the expression of GFP was observed. It was verified that the introduction efficiency was nearly 100% and the GFP gene was expressed at very high levels in all the cells (Figure 15).

<Introduction into mouse primary bone marrow cells>

[0144] Further, an experiment was conducted, in which mouse primary bone marrow cells were separated by utilizing lineage markers and were infected with F-deficient SeV vector. First, 5-fluorouracil (5-FU, Wako Pure Chemical Industries) was given to C57BL mouse (6-week old male) at a dose of 150 mg/kg by intraperitoneal injection (IP injection); 2 days after the administration, bone marrow cells were collected from the thighbone. The mononuclear cells were separated by density gradient centrifugation using Lympholyte-M (Cedarlane). A mixture (3x 10⁷) of Streptavidin-magnetic beads (Pharmingen; Funakoshi), which had been coated with biotin-labeled anti-CD45R (B220), anti-Ly6G (Gr-1), anti-Ly-76 (TER-119), anti-1 (Thyl.2), and anti-Mac-1, were added to the mononuclear cells (3x 10⁶ cells), and the resulting mixture was allowed to react at 4°C for 1 hour; a fraction, from which Lin⁺ cells had been removed by a magnet, was recovered (Lin⁻ cells) (Erich, S. et al., Blood 1999. 93 (1), 80-86). SeV of 2x 10⁷ HAU/ml was added to 4x 10⁵ cells of Lin cell, and further recombinant rat SCF (100 ng/ml, BRL) and recombinant human IL-6 (100 U/ml) were added thereto. In addition, F-deficient SeV of 4x 10⁷ HAU/ml was added to 8x 10⁵ of total bone marrow cells, and GFP-SeV of 5x 10⁷ HAU/ml was added to 1x 10⁶ cells. GFP-SeV was prepared by inserting a PCR-amplified NotI fragment, which contains the green fluorescence protein (GFP) gene (the length of the structural gene is 717 bp) to which a transcription initiation (R1), a termination (R2) signal and an intervening (IG) sequence are added, at the restriction enzyme NotI-cleavage site of SeV transcription unit pUC18/T7HVJRz. DNA (+18) (Genes Cells, 1996, 1: 569-579). The reconstitution of viruses comprising the GFP gene was performed according to a known method (Genes Cells, 1996, 1: 569-579), using LLC-MK2 cells and embryonated egg, and then the viruses comprising the gene of interest were recovered. After a 48-hour culture following the infection with GFP-SeV, the cells were divided into two groups; one of them was allowed to react to phycoerythrin(PE)-labeled anti-CD117 (c-kit; Pharmingen) for 1 hour; the other was a control group. The cells were washed 3 times with PBS then were analyzed in a flow cytometer (EPICS Elite ESP; Coulter, Miami, FL).

[0145] The result showed that F-deficient SeV vector was also infected to bone marrow cells enriched by anti-c-kit antibody that has been utilized as a marker for blood primitive stem cells and the expression of the GFP gene was observed (Figure 16). The presence of infective particles in the culture supernatant was confirmed by determining the presence of GFP-expressing cells three days after the addition of cell culture supernatant treated with trypsin to LLC-MK2 cells. It was clarified that none of these cells released infective virus particles.

[Example 6] Vector administration into rat cerebral ventricle

[0146] Rats (F334/Du Crj, 6 week old, female, Charles River) were anesthetized by intraperitoneal injection of Nembutal sodium solution (Dainabot) diluted 10 folds (5 mg/ml) with physiological saline (Otsuka Pharmaceutical Co., Ltd.). Virus was administrated using brain stereotaxic apparatus for small animals (DAVID KOPF). 20 µl (10⁸ CIU) were injected at the point 5.2 mm toward bregma from interaural line, 2.0 mm toward right ear from lambda, 2.4 mm beneath the brain surface, using 30G exchangeable needles (Hamilton). A high level expression of GFP protein was observed in ventricle ependymal cells (Figure 17). Furthermore, in the case of F deficient SeV vector, the expression of GFP protein was observed only in ependymal cells or nerve cells around the injection site, which come into contact with the virus, and no lesion was found in this region. Abnormality in behavior or changes in body weight were not observed in the administered rats until dissection. After dissection, no lesion was found in the brain or in any of the tissues and organs analyzed, such as liver, lung, kidney, heart, spleen, stomach, intestine, and so forth.

[Example 7] Formation of F-less virus particles from F deficient SeV genome

<1>

[0147] F non-expressing LLC-MK2 cells and F expressing LLC-MK2 cells (LC-MK2/F7) were infected with F deficient SeV virus and cultured with (+) and without (-) trypsin. The result of HA assay of cell culture supernatant after 3 days is shown in Figure 18A. The culture supernatants were inoculated to embryonated chicken eggs, and the result of HA assay of chorioallantoic fluids after a 2 day-culture is shown in Figure 18B. "C" on top of panel indicates PBS used as the control group. The numbers indicated under "Dilution" indicates the dilution fold of the virus solution. Further, HA-positive chorioallantoic fluids in embryonated chicken eggs (lanes 11 and 12) was reinoculated into embryonated chick-

en eggs, and after culturing for two days, the chorioallantoic fluid was examined with HA assay (Figure 19C). As a result, F non-expressing cells or embryonated chicken eggs infected with F deficient SeV virus were found to be HA-positive. However, viruses had not propagated after re-inoculation to embryonated chicken eggs, proving that the HA-positive virus solution does not have secondary infectivity.

<2>

[0148] The non-infectious virus solution amplified in F non-expressing cells was examined for the existence of virus particles. Northern blot analysis was performed for total RNA prepared from the culture supernatant of F expressing cells, HA-positive, non-infectious chorioallantoic fluid, and wildtype SeV by QIAamp viral RNA mini kit (QIAGEN), using the F gene and HN gene as probes. As a result, bands were detected for RNA derived from chorioallantoic fluid or virus in culture supernatant of F expressing cells when the HN gene was used as the probe, whereas no bands were detected when using the F gene probe (Figure 10). It was proven that the HA-positive, non-infectious fluid has non-infectious virus-like particles with an F deficient genome. Further, analysis of the HA-positive, non-infectious virus solution by an immunoelectron microscopy revealed the existence of virus particles, and the envelope of virion reacted to the antibody recognizing gold colloid-labeled HN protein, but not to the antibody recognizing gold colloid-labeled F protein (Figure 20). This result showed the existence of F-less virions, proving that the virus can be formed as a virion with HN protein alone, even without the existence of the F protein. It has been shown that SeV virion can form with F alone (Leyer, S. et al., J Gen. Virol 79, 683-687 (1998)), and the present result proved for the first time that SeV virion can be formed with HN protein alone. Thus, the fact that F-less virions can be transiently produced in bulk in embryonated chicken eggs shows that virions packaging SeV F deficient RNP can be produced in bulk.

<3>

[0149] As described above, F-less virus virions transiently amplified in embryonated chicken eggs are not at all infective towards cells infected by the Sendai virus. To confirm that functional RNP structures are packaged in envelopes, F expressing cells and non-expressing cells were, mixed with cationic liposome (DOSPER, Boehringer Mannheim) and transfected by incubation for 15 minutes at room temperature. As a result, GFP-expressing cells were not observed at all when the cells are not mixed with the cationic liposome, whereas all cells expressed GFP when mixed with cationic liposome. In F non-expressing cells, GFP expression was seen only in individual cells and did not extend to adjacent cells, whereas in F expressing cells, GFP-expressing cells extended to form colonies (Figure Z1). Therefore, it became clear that non-infectious virions transiently amplified in embryonated chicken eggs could express a gene when they are introduced into cells by methods such as transfection.

[Example 8] Reconstitution and amplification of the virus from FHN-deficient SeV genome

<Construction of FHN-deficient genomic cDNA>

[0150] To construct FHN-deficient SeV genomic cDNA (pSeV18⁺/ΔFHN), pUC18/KS was first digested with EcoRI to construct pUC18/Eco, and then whole sequence from start codon of F gene to stop codon of HN gene (4866-8419) was deleted, then it was ligated at BsiwI site (cgtagc). After the sequence of FHN deleted region was confirmed by base sequencing, EcoRI fragment (4057 bp) was recovered from gels to substitute for EcoRI fragment of pUC18/KS to accomplish the construction. A KpnI/SphI fragment (14673 bp) comprising the FHN deleted region was recovered from gels to substitute for KpnI/SphI fragment of pSeV18⁺ to obtain plasmid pSeV18⁺/ΔFHN.

[0151] On the other hand, the construction of FHN-deficient SeV cDNA introduced with GFP was accomplished as follows. Sall/XhoI fragment (7842 bp) was recovered from pSeV18⁺/ΔFRN, and cloned into pGEM11z (Promega). The resultant plasmid was named as pGEM11z/SXdFHN. To the FHN-deficient site of pGEM11z/SXdFHN, PCR product with BsiwI sites at both ends of ATG-TAA (846 bp) of d2EGFP (Clontech) was ligated by digesting with BsiwI enzyme. The resultant plasmid was named as pSeV18⁺/ΔFHN-d2GFP.

<Establishment of FHN-deficient, protein co-expressing cell line>

[0152] The plasmid expressing F gene is identical to the one used for establishment of F deficient, protein co-expressing cell line, and plasmid expressing HN gene was similarly constructed, and the fragment comprising ORF of HN was inserted to unique SwaI site of pCALNdlw (Arai et al., described above) to obtain plasmid named pCALNdlw/HN.

[0153] LLC-MK2 cells were mixed with same amount or different ratio of pCALNdlw/F and pCALNdlw/HN, to introduce genes using mammalian transfection kit (Stratagene), according to the manufacture's protocol. Cells were cloned

after a three week-selection with G418. Drug resistant clones obtained were infected with a recombinant adenovirus (Ade/Cre, Saito et al., described above) (moi=10), which expresses Cre DNA recombinase. Then the cells were collected 3 days after inducing expression of F and HN protein after washing 3 times with PBS(-), and they were probed with monoclonal IgG of anti-SeV F protein and anti-SeV HN protein by using Western blotting method (Figure 22).

<Construction of pGEM/FHN>

[0154] F and HN fragments used for the construction of pCALNdLw/F and pCALNdLw/HN were cloned into pGEM4Z and pGEM3Z (Promega) to obtain pGEM4Z/F and pGEM3Z/HN, respectively. A fragment obtained by PvuII digestion of the region comprising T7 promoter and HN of pGEM3Z/HN was recovered, and ligated into the blunted site cut at the SacI unique site at the downstream of F gene of pGEM4Z/F. F and HN proteins were confirmed by Western blotting using anti-F or anti-HN monoclonal antibodies to be expressed simultaneously when they were aligned in the same direction.

<Reconstitution of FHN-deficient virus>

[0155] The reconstitution of FHN-deficient viruses (P0) was done in two ways. One was using the RNP transfection method as used in the reconstitution of F deficient virus, and the other was using T7 to supply co-expressing plasmids. Namely, under the regulation of T7 promoter, plasmids expressing F and HN proteins were constructed separately, and using those plasmids F and HN proteins were supplied for the reconstitution. In both methods, reconstituted viruses were amplified by FHN coexpressing cells. FHN-deficient, GFP-expressing SeV cDNA (pSeV18*/ΔFHN-d2GFP), pGEM/NP, pGEM/P, pGEM/L, and pGEM/FHN were mixed in the ratio of 12 μg/10 cm dish, 4 μg/10 cm dish, 2 μg/10 cm dish, 4 μg/10 cm dish, and 4 μg/10 cm dish (final total volume, 3 ml/10 cm dish) for gene introduction into LLC-MK2 cells in the same way as F deficient SeV reconstitution described above. Three hours after the gene introduction, media was changed to MEM containing AraC (40 μg/ml, SIGMA) and trypsin (7.5 μg/ml, GIBCO), and cultured further for 3 days. Observation was carried out by fluorescence stereoscopic microscope 2 days after gene introduction. The effect of pGEM/FHN addition was analyzed, and the virus formation was confirmed by the spread of GFP-expressing cells. As a result, a spread of GFP-expressing cells was observed when pGEM/FHN was added at reconstitution, whereas the spread was not observed when pGEM/FHN was not added, and the GFP expression was observed only in a single cell (Figure 23). It is demonstrated that the addition at FHN protein reconstitution caused virus virion formation. On the other hand, in the case of RNP transfection, virus recovery was successfully accomplished in FHN expressing cells of P1, as in the case of F deficiency (Figure 24, upper panel).

[0156] Virus amplification was confirmed after infection of FHN-deficient virus solution to cells induced to express FHN protein 6 hours or more after Ade/Cre infection (Figure 24, lower panel).

[0157] Solution of viruses reconstituted from FHN-deficient GFP-expressing SeV CDNA was infected to LLC-MK2, LLC-MK2/F, LLC-MK2/HN and LLC-MK2/FHN cells, and cultured with or without the addition of trypsin. After 3 days of culture, spread of GFP protein expressing cells was analyzed. As a result, spread of GFP was observed only in LLC-MK2/FHN, confirming that the virus solution can be amplified specifically by FHN co-expression and in a trypsin dependent manner (Figure 25).

[0158] To confirm FHN-deficient viral genome, culture supernatant recovered from LLC-MK2/FHN cells was centrifuged, and RNA was extracted using QIAamp Viral RNA mini kit (QIAGEN), according to manufacturer's protocol. The RNA was used for template synthesis of RT-PCR using Superscript Preamplification System for first Strand Synthesis (GIBCO BRL), and PCR was performed using TAKARA Z-Taq (Takara). F-deficient virus was used as a control group. PCR primer sets were selected as combination of M gene and GFP gene, or combination of M gene and L gene (for combination of M gene and GFP gene (M-GFP), forward: 5'-atcagagacctgacgaatgc / SEQ ID NO: 13, reverse: 5'-aagtcgtgctgctcatgtgg / SEQ ID NO: 14; for combination of M gene and L gene (M-L), forward: 5'-gaaaaacttagggataaagtccc / SEQ ID NO: 15, reverse: 5'-gttatctccggatggtgc / SEQ ID NO: 16). As a result, specific bands were obtained for both F-deficient and FHN-deficient viruses at RT conditions when using M and GFP genes as primers. In the case of using M and L genes as primers, the bands with given size comprising GFP were detected for FHN deficient sample, and lengthened bands with the size comprising HN gene were detected for F deficient one. Thus, FHN deficiency in genome structure was proven (Figure 26).

[0159] On the other hand, FHN-deficient virus was infected to F expressing cells similarly as when using the F-deficient virus, and culture supernatant was recovered after 4 days to perform infection experiment toward LLC-MK2, LLC-MK2/F, and LLC-MK2/FHN. As a result, GFP expression cell was not observed in any infected cell, showing that the virus has no infectiousness to these cells. However, it has been already reported that F protein alone is enough to form virus particles (Kato, A. et al., Genes cells 1, 569-579 (1996)) and that asialoglycoprotein receptor (ASG-R) mediates specific infection to hepatocytes (Spiegel et al., J. Virol 72, 5296-5302, 1998). Thus, virions comprising FHN-deficient RNA genome, with virus envelope configured with only F protein may be released to culture supernatant of

F expressing cells. Therefore, culture supernatant of F expressing cells infected with FHN-deficient virus was recovered, and after centrifugation, RNA was extracted as described above and analyzed by RT-PCR by the method described above. As a result, the existence of RNA comprising FHN-deficient genome was proved as shown in Figure 27.

[0160] Western blotting analysis of virus virion turned into pseudotype with VSV-G clearly shows that F and HN proteins are not expressed. It could be said that herein, the production system of FHN-deficient virus virions was established.

[0161] Moreover, virions released from F protein expressing cells were overlaid on FHN expressing or non-expressing LLC-MK2 cells with or without mixing with a cationic liposome (50 μ l DOSPER/500 μ l/well). As a result, spread of GFP-expressing cells was observed when overlaid as mixture with DOSPER, while HN-less virion only has no infectiousness at all, not showing GFP-expressing cells, as was seen in the case of F-less particles described above. In FHN non-expressing cells GFP expressing cell was observed, but no evidence of virus re-formation and spread was found.

[0162] These virus-like particles recovered from F expressing cells can infect cells continuously expressing ASG-R gene, ASG-R non-expressing cells, or hepatocytes, and whether the infection is liver-specific or ASG-R specific can be examined by the the method of Spiegel et al.

[Example 9] Application of deficient genome RNA virus vector

[0163]

1. F-deficient RNP amplified in the system described above is enclosed by the F-less virus envelope. The envelope can be introduced into cells by adding any desired cell-introducing capability to the envelope by chemical modification methods and such, or by gene introducing reagents or gene guns or the like (RNP transfection, or RNP injection), and the recombinant RNA genome can replicate and produce proteins autonomously and continuously in the cells.

2. A vector capable of specific targeting can be produced, when intracellular domain of HN is left as-is, and the extracellular domain of EN is fused with ligands capable of targeting other receptors in a specific manner, and recombinant gene capable of producing chimeric protein is incorporated into viral-genome. In addition, the vector can be prepared in cells producing the recombinant protein. These vectors can be applicable to gene therapy, as vaccines, or such.

3. Since the reconstitution of SeV virus deficient in both FHN has been successfully accomplished, targeting vector can be produced by introducing targeting-capable envelope chimeric protein gene into FHN deletion site instead of the GFP gene, reconstituting it by the same method as in the case of FHN-deficient vector, amplifying the resultant once in FHN-expressing cells, infecting the resultant to non-expressing cells, and recovering virions formed with only the targeting-capable chimeric envelope protein transcribed from the viral-genome.

4. A mini-genome of Sendai virus and a virion formed with only F protein packaging mini-genome by introducing NP, P, L and F gene to cells have been reported (Leyer et al., J Gen. Virol 79,683-687, 1998). A vector in which murine leukemia virus is turned into pseudo-type by Sendai F protein has also been reported (Spiegel et al., J. Virol 72, 5296-5302, 1998). Also reported so far is the specific targeting of trypsin-cleaved F-protein to hepatocytes mediated by ASG-R (Bitzer et al., J. Virol. 71, 5481-5486, 1997). The systems in former reports are transient particle-forming systems, which make it difficult to continuously recover vector particles. Although Spiegel et al., has reported retrovirus vector turned into pseudo-type by Sendai F protein, this method carries intrinsic problems like the retrovirus being able to introduce genes to only mitotic cells. The virus particles recovered in the present invention with a FHN co-deficient SeV viral-genome and only the F protein as the envelope protein are efficient RNA vectors capable of autonomous replication in the cytoplasm irrespective of cell mitosis. They are novel virus particles, and is a practical system facilitating mass production.

[Example 10] Virus reconstitution and amplification from FHN-deficient SeV genome

[0164] The techniques of reconstitution of infectious virus particles from cDNA that cloned the viral genome has been established for many single strand minus strand RNA viruses such as the Sendai virus, measles virus.

[0165] In most of the systems, reconstitution is carried out by introducing plasmids introduced with cDNA, NP, P, and T, genes at the downstream of T7 promoter into cells and expressing cDNA and each gene using T7 polymerase. To supply T7 polymerase, recombinant vaccinia virus expressing T7 polymerase is mainly used.

[0166] T7 expressing vaccinia virus can express T7 polymerase efficiently in most cells. Although, because of vaccinia virus-induced cytotoxicity, infected cells can live for only 2 or 3 days. In most cases, rifampicin is used as an anti-vaccinia reagent. In the system of Kato et al. (Kato, A. et al., Genes cells 1, 569-579 (1996)), AraC was used together with rifampicin for inhibiting vaccinia virus growth to a minimum level, and efficient reconstitution of Sendai virus.

[0167] However, the reconstitution efficiency of minus strand RNA virus represented by Sendai virus is several particles or less in 1×10^5 cells, far lower than other viruses such as retroviruses. Cytotoxicity due to the vaccinia virus and the complex reconstitution process (transcribed and translated protein separately attaches to bare RNA to form RNP-like structure, and after that, transcription and translation occurs by a polymerase) can be given as reasons for this low reconstitution efficiency.

[0168] In addition to the vaccinia virus, an adeno virus system was examined as a means for supplying T7 polymerase, but no good result was obtained. Vaccinia virus encodes RNA capping enzyme functioning in cytoplasm as the enzyme of itself in addition to T7 polymerase and it is thought that the enzyme enhances the translational efficiency by capping the RNA transcribed by T7 promoter in the cytoplasm. The present invention tried to enhance the reconstitution efficiency of Sendai virus by treating vaccinia virus with Psoralen-Long-wave-UV method to avoid cytotoxicity due to the vaccinia virus.

[0169] By DNA cross-linking with Psoralen and long-wave ultraviolet light, the state in which the replication of virus with DNA genome is inhibited, without effecting early gene expression in particular, can be obtained. The notable effect seen by inactivation of the virus in the system may be attributed to that vaccinia virus having a long genome (Tsung, K. et al., J Virol 70, 165-171 (1996)).

[0170] In the case of wildtype virus that can propagate autonomously, even a single particle of virus formed by reconstitution makes it possible for Sendai virus to be propagated by inoculating transfected cells to embryonated chicken eggs. Therefore, one does not have to consider of the efficiency of reconstitution and the residual vaccinia virus seriously.

[0171] However, in the case of reconstitution of various mutant viruses for researching viral replication, particle formation mechanism, and so on, one may be obligated to use cell lines expressing a protein derived from virus and such, not embryonated chicken eggs, for propagation of the virus. Further, it may greatly possible that the mutant virus or deficient virus propagates markedly slower than the wild type virus.

[0172] To propagate Sendai virus with such mutations, transfected cells should be overlaid onto cells of the next generation and cultured for a long period. In such cases, the reconstitution efficiency and residual titer of vaccinia virus may be problematic. In the present method, titer of surviving vaccinia virus was successfully decreased while increasing reconstitution efficiency.

[0173] Using the present method, a mutant virus that could have not been ever obtained in the former system using a non-treated vaccinia virus was successfully obtained by reconstitution (F, FHN-deficient virus). The present system would be a great tool for the reconstitution of a mutant virus, which would be done more in the future. Therefore, the present inventors examined the amount of Psoralen and ultraviolet light (UV), and the conditions of vaccinia virus inactivation.

<Experiment>

[0174] First, Psoralen concentration was tested with a fixed irradiation time of 2 min. Inactivation was tested by measuring the titer of vaccinia virus by plaque formation, and by measuring T7 polymerase activity by pGEM-luc plasmid under the control of T7 promoter and mini-genome of Sendai virus. The measurement of T7 polymerase activity of mini-genome of Sendai virus is a system in which cells are transfected concomitantly with plasmid of mini-genome of Sendai virus and pGEM/NP, pGEM/P, and pGEM/L plasmids, which express NP-, P-, and L-protein of Sendai virus by T7, to examine transcription of luciferase enzyme protein by RNA polymerase of Sendai virus after the formation of ribonucleoprotein complex.

[0175] After the 2 min UV irradiation, decrease in titer of vaccinia virus depending on psoralen concentration was seen. However, T7 polymerase activity was unchanged for a Psoralen concentration up to 0, 0.3, and 1 $\mu\text{g/ml}$, but decreased approximately to one tenth at 10 $\mu\text{g/ml}$ (Figure 28).

[0176] Furthermore, by fixing Psoralen concentration to 0.3 $\mu\text{g/ml}$, UV irradiation time was examined. In accordance with the increase of irradiation time, the titer of vaccinia virus was decreased, although no effect on T7 polymerase activity was found up to a 30 min irradiation. In this case, under the conditions of 0.3 $\mu\text{g/ml}$ and 30 min irradiation, titer could be decreased down to 1/1000 without affecting T7 polymerase activity (Figure 29).

[0177] However, in vaccinia virus with a decreased titer of 1/1000, CPE 24 hours after infection at $\text{moi}=2$ calibrated to pretreatment titer ($\text{moi}=0.002$ as residual titer after treatment) was not different from that of non-treated virus infected at $\text{moi}=2$ (Figure 30).

[0178] Using vaccinia virus treated under the conditions described above, the efficiency of reconstitution of Sendai virus was examined. Reconstitution was carried out by the procedure described below, modifying the method of Kato et al, mentioned above. LLC-MK2 cells were seeded onto 6-well microplates at 3×10^5 cells/well, and after an overnight culture, vaccinia virus was diluted to the titer of 6×10^5 pfu/100 μl calibrated before PLWUV treatment, and infected to PBS-washed cells. One hour after infection, 100 μl of OPTI-MEM added with 1, 0.5, 1, and 4 μg of plasmid pGEM-NP, P, L, and cDNA, respectively, was further added with 10 μl Superfect (QIAGEN) and left standing for 15 min at room

temperature, and after adding 1 ml OPTI-MEM (GIBCO) (containing Rif. and AraC), was overlaid onto the cells.

[0179] Two, three and four days after transfection, cells were recovered, centrifuged, and suspended in 300 μ l/well of PBS. 100 μ l of cell containing solution made from the suspension itself, or by diluting the suspension by 10 or 100 folds, was inoculated to embryonated chicken eggs at day 10 following fertilization, 4 eggs for each dilution (1×10^5 , 1×10^4 , and 1×10^3 cells, respectively). After 3 days, allantoic fluid was recovered from the eggs and the reconstitution of virus was examined by HA test (Table 1). Eggs with HA activity was scored as 1 point, 10 points and 100 points for eggs inoculated with 1×10^5 , 1×10^4 , and 1×10^3 cells, respectively, to calculate the Reconstitution Score (Figure 31). The formula is as shown in Table 1.

Table 1. Effect of the duration of UV treatment of vaccinia virus on reconstitution efficiency of Sendai virus

The number of inoculated cells	Score (a)	The number of HA positive eggs (b)											
		2d						3d					
		0'			15'			0'			15'		
		1	2	4	4	4	4	0	2	4	0	1	3
10^5	1 (a1)	1	2	4	4	4	4	0	2	4	0	1	3
10^4	10 (a2)	0	1	3	2	2	3	0	2	3	4	0	4
10^3	100 (a3)	0	0	0	1	0	0	0	1	0	0	0	0
Reconstitution Score	(a1+a2+a3) x b	1	12	24	124	0	122	0	122	34	244	1	44

$$\text{Reconstitution Score} = (a1+a2+a3) \times b$$

[0180] Also, residual titers of vaccinia virus measured at 2, 3, and 4 days after transfection within cells were smaller in the treated group in proportion to the titer given before transfection (Figure 32).

[0181] By inactivating vaccinia virus by PLWUV, titer could be decreased down to 1/1000 without affecting T7 polymerase activity. However, CPE derived from vaccinia virus did not differ from that of non-treated virus with a 1000 fold higher titer as revealed by microscopic observations.

[0182] Using vaccinia virus treated with the condition described above for reconstitution of Sendai virus, reconstitution efficiency increased from ten to hundred folds (Figure 31). At the same time, residual titer of vaccinia virus after transfection was not 5 pfu/ 10^5 cells or more. Thus, the survival of replicable vaccinia virus was kept at 0.005% or less.

[Example 11] Construction of pseudotype Sendai virus

<1> Preparation of helper cells in which VSV-G gene product is induced

- 5 **[0183]** Because VSV-G gene product has a cytotoxicity, stable transformant was created in LLC-MK2 cells using plasmid pCALNDLG (Arai T. et al., J. Virology 72 (1998) p1115-1121) in which VSV-G gene product can be induced by Cre recombinase. Introduction of plasmid into LLC-MK2 cells was accomplished by calcium phosphate method (Cal-PhosTMMammalian Transfection Kit, Clontech), according to accompanying manual.
- 10 **[0184]** Ten micrograms of plasmid pCALNDLG was introduced into LLC-MK2 cells grown to 60% confluency in a 10 cm culture dish. Cells were cultured for 24 hours with 10 ml MEM-FCS 10% medium in a 5% CO₂ incubator at 37°C. After 24 hours, cells were scraped off and suspended in 10 ml of medium, and then using five 10 cm culture dishes, 1, 2 and 2 dishes were seeded with 5 ml, 2 ml and 0.5 ml, respectively. Then they were cultured for 14 days in 10 ml MEM-FCS 10% medium containing 1200 µg/ml G418 (GIBCO-BRL) with a medium change on every other day to select stable transformants. Twenty-eight clones resistant to G418 grown in the culture were recovered using cloning rings.
- 15 Each clone was expanded to confluency in a 10 cm culture dish.
- [0185]** For each clone, the expression of VSV-G was examined by Western blotting described below using anti-VSV-G monoclonal antibody, after infection with recombinant adenovirus AxCANCre containing Cre recombinase.
- 20 **[0186]** Each clone was grown in a 6 cm culture dish to confluency, and after that, adenovirus AxCANCre was infected at MOI=10 by the method of Saito et al. (see above), and cultured for 3 days. After removing the culture supernatant, the cells were washed with PBS, and detached from the culture dish by adding 0.5 ml PBS containing 0.05% trypsin and 0.02% EDTA (ethylenediaminetetraacetic acid) and incubating at 37°C, 5 min. After suspending in 3 ml PBS, the cells were collected by centrifugation at 1500x g, 5 min. The cells obtained were resuspended in 2 ml PBS, and then centrifuged again at 1500x g, 5 min to collect cells.
- 25 **[0187]** The cells can be stored at -20°C, and can be used by thawing according to needs. The collected cells were lysed in 100 µl cell lysis solution (RIPA buffer, Boehringer Mannheim), and using whole protein of the cells (1x 10⁵ cells per lane) Western blotting was performed. Cell lysates were dissolved in SDS-polyacrylamide gel electrophoresis sample buffer (buffer comprising 6 mM Tris-HCl (pH6.8), 2% SDS, 10% glycerol, 5% 2-mercaptoethanol) and subjected as samples for electrophoresis after heating at 95°C, 5 min. The samples were separated by electrophoresis using SDS-polyacrylamide gel (Multigel 10/20, Daiichi Pure Chemicals Co., Ltd), and the separated protein was then transferred to transfer membrane (Immobilon-P TransferMembranes, Millipore) by semi-dry blotting method. Transfer was carried out using transfer membrane soaked with 100% methanol for 20 sec and with water for 1 hour, at a 1 mA/cm² constant current for 1 hour.
- 30 **[0188]** The transfer membrane was shaken in 40 ml of blocking solution (Block-Ace, Snow Brand Milk Products Co., Ltd.) for 1 hour, and washed once in PBS.
- 35 **[0189]** The transfer membrane and 5 ml anti-VSV-G antibody (clone P4D4, Sigma) diluted 1/1000 by PBS containing 10% blocking solution were sealed in a vinyl-bag and left to stand at 4°C.
- [0190]** The transfer membrane was soaked twice in 40 ml of PBS-0.1% Tween 20 for 5 min, and after the washing, soaked in PBS for 5 min for washing.
- 40 **[0191]** The transfer membrane and 5 ml of anti-mouse IgG antibody labeled with peroxidase (anti-mouse immunoglobulin, Amersham) diluted to 1/2500 in PBS containing 10% blocking solution were sealed in vinyl-bag and were shaken at room temperature for 1 hour.
- [0192]** After shaking, the transfer membrane was soaked twice in PBS-0.1% Tween 20 for 5 min, and after the washing, soaked in PBS for 5 min for washing.
- 45 **[0193]** The detection of proteins on the membrane crossreacting with anti-VSV-G antibody was carried out by the luminescence method (ECL Western blotting detection reagents, Amersham). The result is shown in Figure 33. Three clones showed AxCANCre infection specific VSV-G expression, confirming the establishment of LLC-MK2 cells in which VSV-G gene product can be induced.
- [0194]** One clone among the clones obtained, named as LLCG-L1, was subjected to flow cytometry analysis using anti-VSV antibody (Figure 34). As a result, reactivity with antibody specific to VSV-G gene induction was detected in LLCG-L1, confirming that VSV-G protein is expressed on the cell surface.
- 50

<2> Preparation of pseudotype Sendai virus comprising a genome deficient in the F gene using helper cells

- 55 **[0195]** Sendai virus comprising a genome deficient in F gene was infected to VSV-G gene expressing cells, and whether production of pseudotype virus with VSV-G as capsid can be seen or not was examined using F-deficient Sendai virus comprising GFP gene described in the examples above, and the expression of GFP gene as an index. As a result, in LLCG-L1 without infection of recombinant adenovirus AxCANCre comprising Cre recombinase, viral gene was introduced by F-deficient Sendai virus infection and GFP-expressing cells were detected, although the

number of expressing cells was not increased. In VSV-G induced cells, chronological increase of GFP-expressing cells was found. When 1/5 of supernatants were further added to newly VSV-G induced cells, no gene introduction was seen in the former supernatant, while the increase of GFP-expressing cells as well as gene introduction were found in the latter supernatant. Also, in the case that supernatant from latter is added to LLC-G-L1 cells without induction of VSV-G, gene was introduced, but increase of GFP-expressing cells was not seen. Taken together, virus propagation specific to VSV-G expressing cells was found, and pseudotype F-deficient virus formation with VSV-G was found.

<3> Evaluation of conditions for producing pseudotype Sendai virus with F gene-deficient genome

[0196] A certain amount of pseudotype Sendai viruses with F gene-deficient genomes was infected changing the amount of AxCANCre infection (MOI=0, 1.25, 2.5, 5, and 10) and culture supernatant was recovered at day 7 or day 8. Then, the supernatant was infected to the cells before and after induction of VSV-G, and after 5 days, number of cells expressing GFP was compared to see the effect of amount of VSV-G gene expression. As a result, no virus production was found at MOI=0 and maximum production was found at MOI=10 (Figure 35). In addition, when time course of virus production was analyzed, the production level started to increase from day 5 or after, persisting to day 8 (Figure 36). The measurement of virus titer was accomplished by calculating the number of particles infected to cells in the virus solution (CIU), by counting GFP-expressing cells 5 days after infection of serially (10 fold each) diluted virus solutions to cells not yet induced with VSV-G. As a result, the maximal virus production was found to be 5×10^5 CIU/ml.

<4> Effect of anti-VSV antibody on infectiousness of pseudotype Sendai virus with F gene-deficient genome

[0197] As to whether pseudotype Sendai virus with F gene-deficient genome obtained by using VSV-G expressing cells comprises VSV-G protein in the capsid, the neutralizing activity of whether infectiousness will be affected was evaluated using anti-VSV antibody. Virus solution and antibody were mixed and left standing at room temperature for 30 min, and then infected to LLC-G-L1 cells without VSV-G induction. On day 5, gene-introducing capability was examined by the existence of GFP-expressing cells. As a result, perfect inhibition of infectiousness was seen by the anti-VSV antibody, whereas in Sendai virus with F gene-deficient genome having the original capsid, the inhibition was not seen (Figure 37). Therefore, it was clearly shown that the present virus obtained is a pseudotype Sendai virus comprising VSV-G protein in its capsid, in which infectiousness of the virus can be specifically inhibited by an antibody.

<5> Confirmation of pseudotype Sendai virus's possession of F-deficient genome

[0198] Western blotting analysis of cell extract of infected cells was carried out to examine if the present virus propagated in cells expressing VSV-G gene is the F-deficient type. Western analysis was accomplished by the method described above. As the primary antibodies, anti-Sendai virus polyclonal antibody prepared from rabbit, anti-F protein monoclonal antibody prepared from mouse, and anti-HN protein monoclonal antibody prepared from mouse were used. As the secondary antibodies, anti-rabbit IgG antibody labeled with peroxidase in the case of anti-Sendai virus polyclonal antibody, and anti-mouse IgG antibody labeled with peroxidase in the case of anti-F protein monoclonal antibody and anti-HN protein monoclonal antibody, were used. As a result, F protein was not detected, whereas protein derived from Sendai virus and HN protein were detected, confirming it is F-deficient type.

<6> Preparation of pseudotype Sendai virus with F and HN gene-deficient genome by using helper cells

[0199] Whether the production of pseudotype virus with VSV-G in its capsid is observed after the infection of Sendai virus with F and HN gene-deficient genome to LLC-G-L1 cells expressing VSV-G gene was analyzed using GFP gene expression as the indicator and F and HN gene-deficient Sendai virus comprising GFP gene described in examples above, by a similar method as described in examples above. As a result, virus propagation specific to VSV-G expressing cells was observed, and the production of F and HN deficient Sendai virus that is a pseudotype with VSV-G was observed (Figure 38). The measurement of virus titer was accomplished by calculating the number of particles infected to cells in the virus solution (CIU), by counting GFP-expressing cells 5 days after infection of serially (10 fold each) diluted virus solutions to cells not yet induced with VSV-G. As a result, the maximal virus production was 1×10^6 CIU/ml.

<7> Confirmation of pseudotype Sendai virus's possession of F and HN deficient genome

[0200] Western blotting of proteins in cell extract of infected cells was carried out to analyze whether the present virus propagated in VSV-G expressing cells are the F and HN deficient type. As a result, F and HN proteins were not detected, whereas proteins derived from Sendai virus were detected, confirming that it is F and HN deficient type

(Figure 39).

[Example 12] Analysis of virus reconstitution method

5 <Conventional method>

[0201] LLC-MK2 cells were seeded onto 100 mm culture dishes at 5×10^6 cells/dish. After a 24 hour culture, the cells were washed once with MEM medium without serum, and then infected with recombinant vaccinia virus expressing T7 RNA polymerase (Fuerst, T.R. et al., Proc. Natl. Acad. Sci. USA 83, 8122-8126 1986) (vTF7-3) at room temperature for 1 hour (moi=2) (moi=2 to 3, preferably moi=2 is used). The virus used herein, was pretreated with 3 $\mu\text{g/ml}$ psoralen and long-wave ultraviolet light (365 nm) for 5 min. Plasmids pSeV18 Δ F-GFP, pGEM/NP, pGEM/P, and pGEM/L (Kato, A. et al., Genes cells 1, 569-579(1996)) were suspended in Opti-MEM medium (GIBCO) at ratio of 12 μg , 4 μg , 2 μg , and 4 $\mu\text{g/dish}$, respectively. Then, SuperFect transfection reagent (1 $\mu\text{g DNA/5 } \mu\text{l}$, QIAGEN) was added and left to stand at room temperature for 15 min and 3 ml Opti-MEM medium containing 3% FBS was added. Thereafter, the cells were washed twice with MEM medium without serum, and DNA-SuperFect mixture was added. After a 3 hr culture, cells were washed twice with MEM medium without serum, and cultured 70 hours in MEM medium containing 40 $\mu\text{g/ml}$ cytosine β -D-arabinofuranoside (AraC, Sigma). Cells and culture supernatant were collected as P0-d3 samples. Pellets of P0-d3 were suspended in Opti-MEM medium (10^7 cells/ml). They were freeze-thawed three times and then mixed with lipofection reagent DOSPER (Boehringer Mannheim) (10^6 cells/25 μl DOSPER) and left to stand at room temperature for 15 min. Then, F expressing LLC-MK2/F7 cells were transfected with the mixture (10^6 cells/well in 24-well plate) and cultured with MEM medium without serum (containing 40 $\mu\text{g/ml}$ AraC and 7.5 $\mu\text{g/ml}$ trypsin). Culture supernatants were recovered on day 3 and day 7 and were designated as P1-d3 and P1-d7 samples.

<Envelope plasmid + F expressing cells overlaying method>

[0202] Transfection was carried out similarly as described above, except that 4 $\mu\text{g/dish}$ envelope plasmid pGEM/FHN was added. After a 3 hr culture, cells were washed twice with MEM medium without serum, and cultured 48 hours in MEM medium containing 40 $\mu\text{g/ml}$ cytosine β -D-arabinofuranoside (AraC, Sigma) and 7.5 $\mu\text{g/ml}$ trypsin. After removing the culture supernatant, cells were overlaid with 5 ml cell suspension solution of a 100 mm dish of F expressing LLC-MK2/F7 cells suspended with MEM medium without serum (containing 40 $\mu\text{g/ml}$ AraC and 7.5 $\mu\text{g/ml}$ trypsin). After a 48 hr culture, cells and supernatants were recovered and designated as P0-d4 samples. Pellets of P0-d4 samples were suspended in Opti-MEM medium (2×10^7 cells/ml) and freeze-thawed three times. Then F expressing LLC-MK2/F7 cells were overlaid with the suspension (2×10^6 cells/well, 24-well plate) and cultured in MEM medium without serum (containing 40 $\mu\text{g/ml}$ AraC and 7.5 $\mu\text{g/ml}$ trypsin). Culture supernatants were recovered on day 3 and day 7 of the culture, designated as P1-d3 and P1-d7 samples, respectively. As a control, experiment was carried out using the same method as described above, but without overlaying and adding only the envelope plasmid.

<CIU (Cell Infectious Units) measurement by counting GFP-expressing cells (GFP-CIU)>

[0203] LLC-MK2 cells were seeded onto a 12-well plate at 2×10^5 cells/well, and after 24 hr culture the wells were washed once with MEM medium without serum. Then, the cells were infected with 100 $\mu\text{l/well}$ of appropriately diluted samples described above (P0-d3 or P0-d4, P1-d3, and P1-d7), in which the samples were diluted as containing 10 to 100 positive cells in 10 cm^2 . After 15 min, 1 ml/well of serum-free MEM medium was added, and after a further 24 hr culture, cells were observed under fluorescence microscopy to count GFP-expressing cells.

<Measurement of CIU (Cell Infectious Units)>

[0204] LLC-MK2 cells were seeded onto a 12-well plate at 2×10^5 cells/dish and after a 24 hr culture, cells were washed once with MEM medium without serum. Then, the cells were infected with 100 $\mu\text{l/well}$ of samples described above, in which the virus vector contained is designated as SeV/ Δ F-GFP. After 15 min, 1 ml/well of MEM medium without serum was added and cultured for a further 24 hours. After the culture, cells were washed with PBS (-) three times and were dried up by leaving standing at room temperature for approximately 10 min to 15 min. To fix cells, 1 ml/well acetone was added and immediately removed, and then the cells were dried up again by leaving to stand at room temperature for approximately 10 min to 15 min. 300 $\mu\text{l/well}$ of anti-SeV polyclonal antibody (DN-1) prepared from rabbit, 100-fold diluted with PBS (-) was added to cells were and incubated for 45 min at 37°C . Then, they were washed three times with PBS (-) and 300 $\mu\text{l/well}$ of anti-rabbit IgG (H+L) fluorescence-labeled second antibody (AlexaTM568, Molecular Probes), 200-fold diluted with PBS (-) was added and incubated for 45 min at 37°C . After washing with PBS (-) three times, the cells were observed under fluorescence microscopy (Emission: 560 nm, Absorption: 645

nm filters, Leica) to find florescent cells (Figure 40).

[0205] As controls, samples described above (SeV/ Δ F-GFP) were infected at 100 μ l/well, and after 15 min 1 ml/well of MEM without serum was added, and after a 24 hr culture, cells were observed under fluorescence microscopy (Emission: 360 nm, Absorption: 470 nm filters, Leica) to find GFP-expressing cells, without the process after the culture.

[Example 13] Evaluation of the most suitable PLWUV (Psoralen and Long-Wave UV light) treatment conditions for vaccinia virus (vTF7-3) for increasing reconstitution efficiency of deficient-type Sendai virus vector

[0206] LLC-MK2 cells were seeded onto 100 mm culture dishes at 5×10^6 cells/dish, and after a 24 hr culture, the cells were washed once with MEM medium without serum. Then, the cells were infected with recombinant vaccinia virus (vTF7-3)(Fuerst, T.R. et al., Proc. Natl. Acad. Sci. USA 83, 8122-8126(1986)) expressing T7 RNA polymerase at room temperature for 1 hour (moi=2) (moi=2 to 3, preferably moi=2 is used). The virus used herein, was pretreated with 0.3 to 3 μ g/ml psoralen and long-wave ultraviolet light (365 nm) for 2 to 20 min. Plasmids pSeV18* Δ F-GFP, pGEM/NP, pGEM/P, and pGEM/L (Kato, A. et al., Genes cells 1, 569-579 (1996)) were suspended in Opti-MEM medium (GIBCO) at ratio of 12 μ g, 4 μ g, 2 μ g, and 4 μ g/dish, respectively. Then, SuperFect transfection reagent (1 μ g DNA/5 μ l, QIAGEN) was added and left to stand at room temperature for 15 min and 3 ml Opti-MEM medium containing 3% FBS was added. Thereafter, the cells were washed twice with MEM medium without serum, and then DNA-SuperFect mixture was added. After a 3 hr culture, cells were washed twice with MEM medium without serum, and cultured 48 hours in MEM medium containing 40 μ g/ml cytosine β -D-arabinofuranoside (AraC, Sigma). Approximately 1/20 offield of view in 100 mm culture dish was observed by a fluorescence microscope and GFP-expressing cells were counted. To test the inactivation of vaccinia virus (vTF7-3), titer measurement by plaque formation (Yoshiyuki Nagai et al., virus experiment protocols, p291-296, 1995) was carried out.

[0207] Further, fixing the timing of recovery after transfection to day 3, psoralen and UV irradiation time were examined. Using vaccinia virus (vTF7-3) treated with each PLWUV treatment, reconstitution efficiency of Sendai virus was examined. Reconstitution was carried out by modifying the method of Kato et al., namely by the procedure described below. LLC-MK2 cells were seeded onto a 6-well microplate at 5×10^5 cells/well, and after an overnight culture (cells were considered to grow to 1×10^6 cells/well), PBS washed cells were infected with diluted vaccinia virus (VTF7-3) at 2×10^6 pfu/100 μ l calibrated by titer before PLWUV treatment. After a 1 hour infection, 50 μ l of Opti-MEM medium (GIBCO) was added with 1, 0.5, 1, and 4 μ g of plasmid pGEM/NP, pGEM/P, pGEM/L, and additional type SeV cDNA (pSeV18*b (+)) (Hasan, M. K, et al., J. General Virology 78: 2813-2820, 1997), respectively. 10 μ l SuperFect (QIAGEN) was further added and left to stand at room temperature for 15 min. Then, 1 ml of Opti-MEM (containing 40 μ g/ml AraC) was added and overlaid onto the cells. Cells were recovered 3 days after transfection, then centrifuged and suspended in 100 μ l/well PBS. The suspension was diluted 10, 100, and 1000-fold and 100 μ l of resultant cell solution was inoculated into embryonated chicken eggs 10 days after fertilization, using 3 eggs for each dilution (1×10^5 , 1×10^4 and 1×10^3 cells, respectively). After 3 days, allantoic fluid was recovered from the eggs and virus reconstitution was examined by HA test. To calculate reconstitution efficiency, eggs showing HA activity that were inoculated with 1×10^5 cells, 1×10^4 cells and 1×10^3 cells, were counted as 1, 10, and 100 point(s), respectively.

<Results>

[0208] Results of Examples 12 and 13 are shown in Figures 40 to 43, and Table 2. The combination of envelope expressing plasmid and cell overlay increased the reconstitution efficiency of SeV/ Δ F-GFP. Notable improvement was obtained in d3 to d4 (day 3 to day4) of P0 (before subculture) (Figure 41). In Table 2, eggs were inoculated with cells 3 days after transfection. The highest reconstitution efficiency was obtained in day 3 when treated with 0.3 μ g/ml psoralen for 20 min. Thus, these conditions were taken as optimal conditions (Table 2).

Table 2: Effect of PLWUV treatment of vaccinia virus on reconstitution of Sendai virus

Effect of PLWUV treatment of vaccinia virus on reconstitution of Sendai virus (eggs were inoculated with cells 3 days after transfection)

The number of inoculated cells	Score (a)	The number of HA ⁺ positive eggs (b)							
		0 μ g/ml	0.3 μ g/ml	1 μ g/ml		3 μ g/ml			
		0'	20'	5'	10'	20'	2'	5'	10'
10^5	1(a1)	0	3	3	3	3	3	3	3
10^4	10(a2)	0	3	2	3	3	1	3	1
10^3	100(a3)	0	3	0	1	1	0	1	0
Reconstitution Score	$(a1+a2+a3) \times b$	0	333	43	133	133	13	133	13

Reconstitution Score = (a1+a2+a3) × b

[Example 14] Preparation of LacZ-comprising, F-deficient, GFP-non-comprising Sendai virus vector

<Construction of F-deficient type, LacZ gene-comprising SeV vector cDNA>

[0209] To construct cDNA comprising LacZ gene at Not I restriction site existing at the upstream region of NP gene of pSeV18⁺/ΔF described in Example 1 (pSeV (+18:LacZ)/ΔF), PCR was performed to amplify the LacZ gene. PCR was carried out by adjusting LacZ gene to multiples of 6 (Hausmann, S et al., RNA 2, 1033-1045 (1996)) and using primer (5'-GCGCGGCCGCGTACGGTGGCAACCATGTCGTTTACTTTGACCAA-3' /SEQ ID NO: 17) comprising Not I restriction site for 5' end, and primer (5'-GCGCGGCCGCGATGAACCTTCACCCTAAGTTTTCTTACTACGGCGTACGCTATTACTTC TGACACCAGACCAACTGGTA-3' /SEQ ID NO: 18) comprising transcription termination signal of SeV (E), intervening sequence (I), transcription initiation signal (S), and Not I restriction site for 3' end, using pCMV-β (Clontech) as template. The reaction conditions were as follows. 50 ng pCMV-β, 200 μM dNTP (Pharmacia Biotech), 100 pM primers, 4 U Vent polymerase (New England-Biolab) were mixed with the accompanying buffer, and 25 reaction temperature cycles of 94°C 30 sec, 50°C 1 min, 72°C 2 min were used. Resultant products were electrophoresed with agarose gel electrophoreses. Then, 3.2 kb fragment was cut out and digested with NotI after purification. pSeV(+18: LacZ)/ΔF was obtained by ligating with NotI digested fragment of pSeV18⁺/ΔF.

<Conventional method>

[0210] LLC-MK2 cells were seeded onto 100 mm culture dish at 5×10^6 cells/dish, and after a 24 hour culture, the cells were washed once with MEM medium without serum. Then, the cells were infected with recombinant vaccinia

virus (vTF7-3)(Fuerst, T.R. et al., Proc. Natl Acad. Sci. USA 83, 8122-8126 (1986)) expressing T7 RNA polymerase at room temperature for 1 hour (moi=2) (moi=2 to 3, preferably moi=2 is used). The virus used herein was pretreated with 3 µg/ml psoralen and long-wave ultraviolet light (365 nm) for 5 min. LacZ comprising, F-deficient type Sendai virus vector cDNA (pSeV(+18:LacZ) ΔF), pGEM/NP, pGEM/P, and pGEM/L (Kato, A. et al., Genes Cells 1, 569-579 (1996)) were suspended in Opti-MEM medium (GIBCO) at a ratio of 12 µg, 4 µg, 2 µg, and 4 µg/dish, respectively, 4 µg/dish envelope plasmid pGEM/FHN and SuperFect transfection reagent (1 µg DNA/5 µl, QIAGEN) were added and left to stand at room temperature for 15 min. Then, 3 ml Opti-MEM medium containing 3% FBS was added and the cells were washed twice with MEM medium without serum, and then the DNA-SuperFect mixture was added. After a 3 hr culture, cells were washed twice with MEM medium without serum, and cultured 24 hours in MEM medium containing 40 µg/ml cytosine β-D-arabinofuranoside (AraC, Sigma) and 7.5 µg/ml trypsin. Culture supernatants were removed and 5 ml of suspension of a 100 mm culture dish of F expressing LLC-MK2/F7 cells in MEM medium without serum (containing 40 µg/ml AraC and 7.5 µg/ml trypsin) was overlaid onto the cells. After further a 48 hr culture, the cells and supernatants were recovered and designated as P0-d3 samples. The P0-d3 pellets were suspended in Opti-MEM medium (2x 10⁷ cells/ml) and after 3 times of freeze-thawing, were mixed with lipofection reagent DOSPER (Boehringer Mannheim) (10⁶ cells/25 µl DOSPER) and left to stand at room temperature for 15 min. Then, F expressing LLC-MK2/F7 cells were transfected with the mixture (10⁶ cells/well, 24-well plate) and cultured with MEM medium without serum (containing 40 µg/ml AraC and 7.5 µg/ml trypsin). The culture supernatants were recovered on day 7, and designated as P1-d7 samples. Further, total volumes of supernatants were infected to F expressing LLC-MK2/F7 cells seeded onto 12-well plates at 37°C for 1 hour. Then, after washing once with MEM medium, the cells were cultured in MEM medium without serum (containing 40 µg/ml AraC and 7.5 µg/ml trypsin). The culture supernatants were recovered on day 7, and were designated as P2-d7 samples. Further, total volumes of supernatants were infected to F expressing LLC-MK2/F7 cells seeded onto 6-well plates at 37°C for 1 hour. Then, after washing once with MEM medium, the cells were cultured in MEM medium without serum (containing 7.5 µg/ml trypsin). The culture supernatants were recovered on day 7, and were designated as P3-d7 samples. Further, total volumes of supernatants were infected to F expressing LLC-MK2/F7 cells seeded onto 10 cm plates at 37°C for 1 hour. Then, after washing once with MEM medium, the cells were cultured in MEM medium without serum (containing 40 µg/ml AraC and 7.5 µg/ml trypsin). The culture supernatants were recovered on day 7, and were designated as P4-d7 samples.

<Measurement of CIU by counting LacZ-expressing cells (LacZ-CIU)>

[0211] LLC-MK2 cells were seeded onto 6-well plate at 2.5x 10⁶ cells/well, and after a 24 hr culture, the cells were washed once with MEM medium without serum and infected with 1/10 fold serial dilution series of P3-d7 made using MEM medium at 37°C for 1 hour. Then, the cells were washed once with MEM medium and 1.5 ml MEM medium containing 10% serum was added. After a three day culture at 37°C, cells were stained with β-Gal staining kit (Invitrogen). Result of experiment repeated three times is shown in Figure 44. As the result of counting LacZ staining positive cell number, 1x 10⁶ CIU/ml virus was obtained in P3-d7 samples in any case.

[Example 15] Regulation of gene expression levels using polarity effect in Sendai virus

<Construction of SeV genomic cDNA>

[0212] Additional NotI sites were introduced into Sendai virus (SeV) full length genomic cDNA, namely pSeV(+) (Kato, A. et al., Genes to Cells 1: 569-579, 1996), in between start signal and ATG translation initiation signal of respective genes. Specifically, fragments of pSeV(+) digested with SphI/SalI (2645 bp), ClaI (3246 bp), and ClaI/EcoRI (5146 bp) were separated with agarose gel electrophoreses and corresponding bands were cut out and then recovered and purified with QIAEXII Gel Extraction System (QIAGEN) as shown in Figure 45(A). The SphI/SalI digested fragment, ClaI digested fragment, and ClaI/EcoRI digested fragment were ligated to LITMUS38 (NEW ENGLAND BIOLABS), pBluescriptII KS+ (STRATAGENE), and pBluescriptII KS+ (STRATAGENE), respectively, for subcloning. Quickchange Site-Directed Mutagenesis kit (STRATAGENE) was used for successive introduction of NotI sites. Primers synthesized and used for each introduction were, sense strand: 5'-ccaccgaccacacccagcgccgacagccacggcttcgg-3' (SEQ ID NO: 19), antisense strand: 5'-ccgaagccgtggtgtcgcggcgcctgggtgtggtcggtgg-3' (SEQ ID NO: 20) for NP-P, sense strand: 5'-gaaatttcacctaagcggcgcaatggcagatatctatag-3' (SEQ ID NO: 21), antisense strand: 5'-ctatagatatctgccattgcccgccttaggtgaaatttc-3' (SEQ ID NO: 22) for P-M, sense strand: 5'-gggataaagtccctgcggccgcttggtgcaaaactctcccc-3' (SEQ ID NO: 23), antisense strand: 5'-ggggagagttttgcaaccaagcggcgcaagggaactttatccc-3' (SEQ ID NO: 24) for M-F, sense strand: 5'-ggtcgcgcggtacttagcggcgccctcaacaagcacagatcatgg-3' (SEQ ID NO: 25), antisense strand: 5'-ccatgatctgtgctgtttgagggcgccgctaagtagcggcgacc-3' (SEQ ID NO: 26) for F-HN, sense strand: 5'-cctgcccattcatgacatagcggcgccgttcccat-tcaccctggg-3' (SEQ ID NO: 27), antisense strand: 5'-cccagggtgaatgggaagcggcgctaggtcatggtgggcagg-3' (SEQ ID NO: 28) for HN-L.

[0213] As templates, Sall/SphI fragment for NP-P, ClaI fragments for P-M and M-F, and ClaI/EcoRI fragments for F-HN and HN-L, which were subcloned as described above were used, and introduction was carried out according to the protocol accompanying Quickchange Site-Directed Mutagenesis kit. Resultants were digested again with the same enzyme used for subcloning, recovered, and purified. Then, they were assembled to Sendai virus genomic cDNA. As a result, 5 kinds of genomic cDNA of Sendai virus (pSeV (+) NPP, pSeV(+)PM, pSeV(+)MF, pSeV(+)FHN, and pSeV (+) HNL) in which NotI sites are introduced between each gene were constructed as shown in Figure 45(B).

[0214] As a reporter gene to test gene expression level, human secreted type alkaline phosphatase (SEAP) was subcloned by PCR. As primers, 5' primer: 5'-gcggcgcgcgcctatctgtctgaagcgccgcgcgcg-3' (SEQ ID NO: 29) and 3' primer: 5'-gcggcgcgcgcctatctgtctgaagcgccgcgcgcg-3' (SEQ ID NO: 30) added with Ascl restriction sites were synthesized and PCR was performed. pSEAP-Basic (CLONTECH) was used as template and Pfu turbo DNA polymerase (STRATAGENE) was used as enzyme. After PCR, resultant products were digested with Ascl, then recovered and purified by electrophoreses, As plasmid for subcloning, pBluescriptII KS+ incorporated in iLs NotI site with synthesized double strand DNA [sense strand: 5'-gcgccgcgcgtttaacgcgcgcgcattaaatccgtagtaagaaaaacttaggtgaaagt tcatcgcg-gccgc-3' (SEQ ID NO: 31), antisense strand: 5'-gcgccgcgcgtgaactttcacctaagttttcttactacggatttaaatggcgccggt taaacgcgcgcgc-3' (SEQ ID NO: 32)] comprising multicloning site (PmeI-Ascl-SwaI) and termination signal-intervening sequence-initiation signal was constructed (Figure 46). To Ascl site of the plasmid, recovered and purified PCR product was ligated and cloned. The resultant was digested with NotI and the SEAP gene fragment was recovered and purified by electrophoreses to ligate into 5 types of Sendai virus genomic cDNA and NotI site of pSeV18+ respectively. The resultant virus vectors were designated as pSeV(+)NPP/SEAP, pSeV(+)PM/SEAP, pSeV(+)MF/SEAP, pSeV(+)FHN/SEAP, pSeV(+)HNL/SEAP, and pSeV18(+)/SEAP, respectively.

<Virus reconstitution>

[0215] LLC-MK2 cells were seeded onto 100 mm culture dishes at 2×10^6 cells/dish, and after 24 hour culture the cells were infected with recombinant vaccinia virus (PLWUV-VacT7) (Fuerst, T.R. et al., Proc. Natl. Acad. Sci. USA 83: 8122-8126, 1986, Kato, A. et al., Genes Cells 1: 569-579, 1996) expressing T7 polymerase for 1 hour (moi=2) at room temperature for 1 hour, in which the virus was pretreated with psoralen and UV. Each Sendai virus cDNA incorporated with SEAP, pGEM/NP, pGEM/P, and pGEM/L were suspended in Opti-MEM medium (GIBCO) at ratio of 12 μ g, 4 μ g, 2 μ g, and 4 μ g/dish, respectively, 110 μ l of SuperFect transfection reagent (QIAGEN) was added, and left to stand at room temperature for 15 min and 3 ml Opti-MEM medium containing 3% FBS was added. Then, the cells were washed and DNA-SuperFect mixture was added. After a 3 to 5 hour culture, cells were washed twice with MEM medium without serum, and cultured 72 hours in MEM medium containing cytosine β -D-arabinofuranoside (AraC). These cells were recovered and the pellets were suspended with 1 ml PBS, freeze-thawed three times. The 100 μ l of resultant was inoculated into chicken eggs, which was preincubated 10 days, and further incubated 3 days at 35°C, then, allantoic fluid was recovered. The recovered allantoic fluids were diluted to 10^{-5} to 10^{-7} and re-inoculated to chicken eggs to make it vaccinia virus-free, then recovered similarly and stocked in aliquots at -80°C. The virus vectors were designated as SeVNPP/SEAP, SeVPM/SEAP, SeVMF/SEAP, SeVFHN/SEAP, SeVHNL/SEAP, and SeV18/SEAP.

<Titer measurement by plaque assay>

[0216] CV-1 cells were seeded onto 6-well plates at 5×10^5 cells/well and cultured for 24 hours. After washing with PBS, cells were incubated 1 hour with recombinant SeV diluted as 10^{-3} , 10^{-4} , 10^{-5} , 10^{-6} and 10^{-7} by BSA/PBS (1% BSA in PBS), washed again with PBS, then overlaid with 3 ml/well of BSA/MEM/agarose (0.2% BSA + 2x MEM, mixed with equivalent volume of 2% agarose) and cultured at 37°C, 0.5% CO₂ for 6 days. After the culture, 3 ml of ethanol/acetic acid (ethanol:acetic acid=1:5) was added and left to stand for 3 hours, then removed with agarose. After washing three times with PBS, cells were incubated with rabbit anti-Sendai virus antibody diluted 100-folds at room temperature for 1 hour. Then, after washing three times with PBS, cells were incubated with Alexa Flour™ labeled goat anti rabbit Ig(G+H)(Molecular Probe) diluted 200-folds at room temperature for 1 hour. After washing three times with PBS, fluorescence images were obtained by lumino-image analyzer LAS1000 (Fuji Film) and plaques were measured. Results are shown in Figure 47. In addition, results of titers obtained are shown in Table 3.

Table 3:

Results of titers of each recombinant Sendai virus measured from results of plaque assay	
Recombinant virus	Titer (pfu/ml)
SeV18/SEAP	3.9×10^9
SeVNPP/SEAP	4.7×10^8

Table 3: (continued)

Recombinant virus	Titer (pfu/ml)
SeVPM/SEAP	3.8X10 ⁹
SeVMF/SEAP	1.5X10 ¹⁰
SeVFHN/SEAP	7.0X10 ⁹
SeVHNL/SEAP	7.1X10 ⁹

<Comparison of reporter gene expression>

[0217] LLC-MK2 cells were seeded onto a 6-well plate at 1 to 5x 10⁵ cells/well and after a 24 hour culture, each virus vector was infected at moi=2. After 24 hours, 100 µl of culture supernatants was recovered and SEAP assay was carried out. Assay was accomplished with Reporter Assay Kit -SEAP- (Toyobo) and measured by lumino-image analyzer LAS1000 (Fuji Film). The measured values were indicated as relative values by designating value of SeV18+/SERF as 100. As a result, SEAP activity was detected regardless of the position SEAP gene was inserted, indicated in Figure 48. SEAP activity was found to decrease towards the downstream of the genome, namely the expression level decreased. In addition, when SEAP gene is inserted in between NP and P genes, an intermediate expression level was detected, in comparison to when SEAP gene is inserted in the upstream of NP gene and when SEAP gene is inserted between P and M genes.

[Example 16] Increase of propagation efficiency of deficient SeV by double deficient ΔF-HN overlay method

[0218] Since the SeV virus reconstitution method used now utilizes a recombinant vaccinia virus expressing T7 RNA polymerase (vTF7-3), a portion of the infected cells is killed by the cytotoxicity of the vaccinia virus. In addition, virus propagation is possible only in a portion of cells and it is preferable if virus propagation could be done efficiently and persistently in a more cells. However, in the case of paramyxovirus, cell fusion occurs when F and HN protein of the same kind virus exists on the cells surface at the same time, causing syncytium formation (Lamb and Kolakofsky, 1996, Fields virology, p1189). Therefore, FHN co-expressing cells were difficult to subculture. Therefore, the inventors thought that recovery efficiency of deficient virus may increase by overlaying helper cells expressing deleted protein (F and HN) to the reconstituted cells. By examining overlaying cells with different times of FHN expression, virus recovery efficiency of FHN co-deficient virus was notably increased.

[0219] LLC-MK2 cells (1x 10⁷ cells/dish) grown to 100% confluency in 10 cm culture dishes was infected with PLWUV-treated vaccinia virus at moi=2 for 1 hour at room temperature. After that, mixing 12 µg/10 cm dish, 4 µg/10 cm dish, 2 µg/10 cm dish, 4 µg/10 cm dish, and 4 µg/10 cm dish of FHN-deficient cDNA comprising d2EGFP (pSeV18+/ΔFHN-d2GFP (Example 8)), pGEM/NP, pGEM/P, pGEM/L, and pGEM/FHN, respectively (3 ml/10cm dish as final volume), and using gene introduction reagent SuperFect (QIAGEN), LLC-MK2 cells were introduced with genes using a method similar to that as described above for the reconstitution of F-deficient virus. After 3 hours, cells were washed three times with medium without serum, then, the detached cells were recovered by slow-speed centrifugation (1000 rpm/2 min) and suspended in serum free MEM medium containing 40 µg/ml AraC (Sigma) and 7.5 µg/ml trypsin (GIBCO) and added to cells and cultured overnight. FHN co-expressing cells separately prepared, which were 100% confluent 10 cm culture dishes, were induced with adenovirus AxCANCre at MOI=10, and cells at 4 hours, 6 hours, 8 hours, day 2, and day 3 were washed once with 5 ml PBS(-) and detached by cell dissociation solution (Sigma). Cells were collected by slow speed centrifugation (1000 rpm/2 min) and suspended in serum free MEM medium containing 40 µg/ml AraC (Sigma) and 7.5 µg/ml trypsin (GIBCO). This was then added to cells in which FHN co-deficient virus was reconstituted (P0) and cultured overnight. Two days after overlaying the cells, cells were observed using fluorescence microscopy to confirm the spread of virus by GFP expression within the cells. The results are shown in Figure 49. When compared to the conventional case (left panel) without overlaying with cells, notably more GFP-expressing cells were observed when cells were overlaid with cells (right). These cells were recovered, suspended with 10⁷ cells/ml of Opti-MEM medium (GIBCO) and freeze-thawed for three times to prepare a cell lysate. Then, FHN co-expressing cells 2 days after induction were infected with the lysate at 10⁶ cells/100 µl/well, and cultured 2 days in serum free MEM medium containing 40 µg/ml AraC (Sigma) and 7.5 µg/ml trypsin (GIBCO) at 37°C in a 5% CO₂ incubator, and the virus titer of culture supernatant of P1 cells were measured by CIU-GFP (Table 4). As a result, no virus amplification effect was detected 4 hours after FHN induction, and notable amplification effects were detected 6 hours or more after induction due to cell overlaying. Especially, viruses released into P1 cell culture supernatant were 10 times more after 6 hours when cell overlaying was done compared to when cell overlaying was not done.

Table 4:

Amplification of deficient SeV by double deficient Δ F-HN cell overlay method					
GFP -CIU	x10 ³ /ml				
	FHNcell*ad/cre				
FHN cell-	4h	6h	8h	2d	3d
8-10	6-9	80-100	70-100	60-100	20-50

[Example 17] Confirmation of pseudotype Sendai virus's possession of F-deficient genome

[0220] Western analysis of proteins of extracts of infected cells was carried out to confirm that the virus propagated by VSV-G gene expression described above is F-deficient type. As a result, proteins derived from Sendai virus were detected, whereas F protein was not detected, confirming that the virus is F-deficient type (Figure 50).

[Example 18] Effect of anti-VSV antibody on infectiousness of pseudotype Sendai virus comprising F and HN gene-deficient genome

[0221] To find out whether pseudotype Sendai virus comprising F and HN gene-deficient genome, which was obtained by using VSV-G expressing line, comprises VSV-G protein in its capsid, neutralizing activity of whether or not infectiousness is affected was examined using anti-VSV antibody. Virus solution and antibody were mixed and left to stand for 30 min at room temperature. Then, LLCG-L1 cells in which VSV-G expression has not been induced were infected with the mixture and gene-introducing capability on day 4 was analyzed by the existence of GFP-expressing cells. As a result, perfect inhibition of infectiousness was seen by anti-VSV antibody in the pseudotype Sendai virus comprising F and HN gene-deficient genome (VSV-G in the Figure), whereas no inhibition was detected in Sendai virus comprising proper capsid (F, HN in the Figure) (Figure 51). Thus, the virus obtained in the present example was proven to be pseudotype Sendai virus comprising VSV-G protein as its capsid, and that its infectiousness can be specifically inhibited by the antibody.

[Example 19] Purification of pseudotype Sendai viruses comprising F gene-deficient and F and HN gene-deficient genomes by density gradient ultracentrifugation

[0222] Using culture supernatant of virus infected cells, sucrose density gradient centrifugation was carried out, to fractionate and purify pseudotype Sendai virus comprising deficient genomes of F gene and F and HN genes. Virus solution was added onto a sucrose solution with a 20 to 60% gradient, then ultracentrifuged for 15 to 16 hours at 29000 rpm using SW41 rotor (Beckman). After ultracentrifugation, a hole was made at the bottom of the tube, then 300 μ l fractions were collected using a fraction collector. For each fraction, Western analysis were carried out to test that the virus is a pseudotype Sendai virus comprising a genome deficient in F gene or F and HN genes, and VSV-G protein as capsid. Western analysis was accomplished by the method as described above. As a result, in F-deficient pseudotype Sendai virus, proteins derived from the Sendai virus, HN protein, and VSV-G protein were detected in the same fraction, whereas F protein was not detected, confirming that it is a F-deficient pseudotype Sendai virus. On the other hand, in F and HN-deficient pseudotype Sendai virus, proteins derived from Sendai virus, and VSV-G protein were detected in the same fraction, whereas F and HN protein was not detected, confirming that it is F and HN deficient pseudotype Sendai virus (Figure 52).

[Example 20] Overcoming of haemagglutination by pseudotype Sendai virus comprising F gene-deficient and F and HN gene-deficient genomes

[0223] LLC-MK2 cells were infected with either pseudotype Sendai virus comprising F gene-deficient or F and HN gene-deficient genome, or Sendai virus with normal capsid, and on day 3, 1% avian red blood cell suspension was added, and left to stand for 30 min at 4°C. Thereafter, cell surface of infected cells expressing GFP were observed. As a result, for virus with F gene-deficient genome and F-deficient pseudotype Sendai virus (SeV/ Δ F, and pseudotype SeV/ Δ F (VSV-G) by VSV-G), agglutination reaction was observed on the surface of infected cells, as well as for the Sendai virus with the original capsid. On the other hand, no agglutination reaction was observed on the surface of infected cells for pseudotype Sendai virus comprising F and HN gene-deficient genome (SeV/ Δ F-HN(VSV-G)) (Figure 53).

[Example 21] Infection specificity of VSV-G pseudotype Sendai virus comprising F gene-deficient genome to cultured cells

[0224] Infection efficiency of VSV-G pseudotype Sendai virus comprising F gene-deficient genome to cultured cells was measured by the degree of GFP expression in surviving cells 3 days after infection using flow cytometry. LLC-MK2 cells showing almost the same infection efficiency in pseudotype Sendai virus comprising F gene-deficient genome and Sendai virus with original capsid were used as controls for comparison. As a result, no difference in infection efficiency was found in human ovary cancer HRA cells, whereas in Jurkat cells of T cell lineage about 2-fold increase in infection efficiency of VSV-G pseudotype Sendai virus comprising F gene-deficient genome was observed compared to controls (Figure 54).

[Example 22] Construction of F-deficient type Sendai virus vector comprising NGF

<Reconstitution of NGF/SeV/ΔF>

[0225] Reconstitution of NGF/SeV/ΔF was accomplished according to the above described "Envelope plasmid + F expressing cells overlaying method". Measurement of titer was accomplished by a method using anti-SeV polyclonal antibody.

<Confirmation of virus genome of NGF/SeV/ΔF (RT-PCR)>

[0226] To confirm NGF/SeV/ΔF virus genome (Figure 55, upper panel), culture supernatant recovered from LLC-MK2/F7 cells were centrifuged, and RNA was extracted using QIAamp Viral RNAmini kit (QIAGEN) according to the manufacturer's protocol. Using the RNA template, synthesis and PCR of RT-PCR was carried out using SUPER-SCRIPT™ ONE-STEP™ RT-PCR SYSTEM (GIBCO BRL). As control groups, additional type SeV cDNA (pSeV18*b (+)) (Hasan, M. K. et al., J. General Virology 78: 2813-2820, 1997) was used. NGF-N and NGF-C were used as PCR primers. For NGF-N, forward: ACTTGCGGCGCCGCAAAGTTCAGTAATGTCCATGTTGTTCTACACTCTG (SEQ ID NO: 33), and for NGF-C, reverse: ATCCGCGGCGCGCATGAACCTTCCACCCTAAGTTTTCTTACTACGGTCAGCCTCT-TCTTGATGC CTTCTG (SEQ ID NO: 34) were used. As a result, when NGF-N and NGF-C were used as primers, an NGF specific band was detected for NGF/SeV/ΔF in the RT conditions. No band was detected for the control group (Figure 55, bottom panel).

[Example 23] NGF protein quantification and measurement of *in vitro* activity expressed after infection of F-deficient type SeV comprising NGF gene

[0227] Infection and NGF protein expression was accomplished using LLC-MK2/F or LLC-MK2 cells grown until almost confluent on culture plates of diameter of 10 cm or 6 cm. NGF/SeV/ΔF and NGF/SeV/ΔF-GFP were infected to LLC-MK2/F cells, and NGF/SeV and GFP/SeV were infected to LLC-MK2 cells at m.o.i 0.01, and cultured 3 days with MEM medium without serum, containing 7.5 μg/ml trypsin (GIBCO). After the 3 day culture, in which almost 100% of cells are infected, medium was changed to MEM medium without trypsin and serum and further cultured for 3 days. Then, each culture supernatant were recovered and centrifuged at 48,000x g for 60 min. Then, quantification of NGF protein and measurement of *in vitro* activity for the supernatant were carried out. Although in the present examples, F-deficient type SeV (NGF/SeV/ΔF, NGF/SeV/ΔF-GFP) (see Figure 55) are infected to LLC-MK2/F cells, if infected with a high m.o.i. (e.g. 1 or 3), namely, infected to cells that are nearly 100% confluent from the beginning, experiment giving similar results can be performed using F non-expressing cells.

[0228] For NGF protein quantification, ELISA kit NGF Emax Immuno Assay System (Promega) and the accompanying protocol were used. 32.4 μg/ml, 37.4 μg/ml, and 10.5 μg/ml of NGF protein were detected in NGF/SeV/ΔF, NGF/SeV/ΔF-GFP, and NGF/SeV infected cell culture supernatant, respectively. In the culture supernatant of NGF/SeV/ΔF and NGF/SeV/ΔF-GFP infected cells, high concentration of NGF protein exists, similar to culture supernatant of NGF/SeV infected cells, confirming that F-deficient type SeV expresses enough NGF.

[0229] The measurement of *in vitro* activity of NGF protein was accomplished by using a dissociated culture of primary chicken dorsal root ganglion (DRG; a sensory neuron of chicken) using survival activity as an index (Nerve Growth Factors (Wiley, New York), pp.95-109 (1989)). Dorsal root ganglion was removed from day 10 chicken embryo, and dispersed after 0.25% trypsin (GIBCO) treatment at 37°C for 20 min. Using high-glucose D-MEM medium containing 100 units/ml penicillin (GIBCO), 100 units/ml streptomycin (GIBCO), 250 ng/ml amphotericin B (GIBCO) 20 μM 2-deoxyuridine (Nakarai), 20 μM 5-fluorodeoxyuridine (Nakarai), 2 mM L-glutamine (Sigma), and 5% serum, cells were seeded onto 96-well plate at about 5000 cells/well. Polylysine precoated 96-well plates (Iwaki) were further coated with laminin (Sigma) before use. At the start point, control NGF protein or previously prepared culture supernatant after

SeV infection was added. After 3 days, cells were observed under a microscope as well as conducting quantification of surviving cells by adding Alamer blue (CosmoBio) and using the reduction activity by mitochondria as an index (measuring 590 nm fluorescence, with 530 nm excitation). Equivalent fluorescence signals were obtained in control (without NGF addition) and where 1/1000 diluted culture supernatant of cells infected with SeV/additional-type-GFP (GFP/SeV) was added, whereas the addition of 1/1000 diluted culture supernatant of cells infected with NGF/SeV/ Δ F, NGF/SeV/ Δ F-GFP, and NGF/SeV caused a notable increase in fluorescence intensity, and was judged as comprising a high number of surviving cells and survival activity (Figure 56). The value of effect was comparable to the addition of amount of NGF protein calculated from ELISA. Observation under a microscope proved a similar effect. Namely, by adding culture supernatant of cells infected with NGF/SeV/ Δ F, NGF/SeV/ Δ F-GFP, and NGF/SeV, increase in surviving cells and notable neurite elongation was observed (Figure 57). Thus, it was confirmed that NGF expressed after infection of NGF-comprising F-deficient type SeV is active form.

[Example 24] Detailed analysis of F-expressing cells

1. moi and induction time course of Adeno-Cre

[0230] By using different moi of Adeno-Cre, LLC-MK2/F cells were infected and after induction of F protein, the amount of protein expression and the change in cell shape were analyzed.

[0231] Expression level was slightly higher in moi=10 compared with moi=1 (Figure 58). When expression amounts were analyzed at time points of 6 h, 12 h, 24 h, and 48 h after induction, high expression level of F protein at 48 h after induction was detected in all cases.

[0232] In addition, changes in cell shape were monitored in a time course as cells were infected with moi=1, 3, 10, 30, and 100. Although a notable difference was found up to moi=10, cytotoxicity was observed for moi=30 or over (Figure 59).

2. Passage number

[0233] After induction of F protein to LLC-MK2/F cells using Adeno-Cre, cells were passaged 7 times and expression level of F protein and the morphology of the cells were analyzed using microscopic observation. On the other hand, laser microscopy was used for analysis of intracellular localization of F protein after induction of F protein in cells passaged until the 20th generation.

[0234] For laser microscopic observation, LLC-MK2/F cells induced with F protein expression were put into the chamber glass and after overnight culture, media were removed and washed once with PBS, then fixed with 3.7% Formalin-PBS for 5 min. Then after washing cells once with PBS, cells were treated with 0.1% Triton X100-PBS for 5 min, and treated with anti-F protein monoclonal antibody (γ -236) (1/100 dilution) and FITC labeled goat anti-rabbit IgG antibody (1/200 dilution) in this order, and finally washed with PBS and observed with a laser microscope.

[0235] As a result, no difference was found in F protein expression levels in cells passaged up to 7 times (Figure 60). No notable difference was observed in morphological change, infectiousness of SeV, and productivity. On the other hand, when cells passaged up to 20 times were analyzed for intracellular localization of F protein using the immune-antibody method, no big difference was found up to 15 passages, but localization tendency of F protein was observed in cells passaged more than 15 times (Figure 61).

[0236] Taken together, cells before 15 passages are considered desirable for the production of F-deficient SeV.

[Example 25] Correlation between GFP-CIU and anti-SeV-CIU

[0237] Correlation of the results of measuring Cell-Infected Unit (CIU) by two methods was analyzed. LLC-MK2 cells were seeded onto a 12-well plate at 2×10^5 cells/dish, and after a 24 hour culture, cells were washed once with MEM medium without serum, and infected with 100 μ l/well SeV/ Δ F-GFP. After 15 min, 1 ml/well serum-free MEM medium was added and further cultured for 24 hours. After the culture, cells were washed three times with PBS(-) and dried up (left to stand for approximately 10 to 15 min at room temperature) and 1 ml/well acetone was added to fix cells and was immediately removed. Cells were dried up again (left to stand for approximately 10 to 15 min at room temperature). Then, 300 μ l/well of anti-SeV polyclonal antibody (DN-1) prepared from rabbits and diluted 1/100 with PBS (-) was added to cells and incubated at 37°C for 45 min and washed three times with PBS (-). Then, to the cells, 300 μ l/well of anti-rabbit IgG(H+D) fluorescence-labeled second antibody (AlexTM 568, Molecular Probes) diluted 1/200 with PBS (-) was added, and incubated at 37°C for 45 min and washed three times with PBS (-). Then, cells with fluorescence were observed under fluorescence microscopy (Emission: 560 nm, Absorption: 645 nm, Filters: Leica).

[0238] As a control, cells were infected with 100 μ l/well of SeV/ Δ F-GFP and after 15 min, 1 ml/well of MEM without serum was added. After a further 24 hour culture, GFP-expressing cells were observed under fluorescence microscopy

(Emission: 360 nm, Absorption: 470 nm, Filters: Leica) without further manipulations.

[0239] A Good correlation was obtained by evaluating the fluorescence intensity by quantification (Figure 62).

[Example 26] Construction of multicloning site

[0240] A multicloning site was added to the SeV vector. The two methods used are listed below.

1. Several restriction sites in full-length genomic cDNA of Sendai virus (SeV) and genomic cDNA of pSeV18⁺ were disrupted, and another restriction site comprising the restriction site disrupted was introduced in between start signal and ATG translation initiation signal of each gene.
2. Into already constructed SeV vector cDNA, multicloning site sequence and transcription initiation signal - intervening sequence-termination signal were added and incorporated into NotI site.

[0241] In the case of method 1, as an introducing method, EagI-digested fragment (2644 bp), ClaI-digested fragment (3246 bp), ClaI/EcoRI-digested fragment (5146 bp), and EcoRI-digested fragment (5010 bp) of pSeV18⁺ were separated by agarose electrophoreses and the corresponding bands were cut out, then it was recovered and purified by QIAEXII Gel Extraction System (QIAGEN). EagI-digested fragment was ligated and subcloned into LITMUS38 (NEW ENGLAND BIOLABS) and ClaI-digested fragment, ClaI/EcoRI-digested fragment, and EcoRI-digested fragment were ligated and subcloned into pBluescriptII KS⁺ (STRATAGENE). Quickchange Site-Directed Mutagenesis kit (STRATAGENE) was used for successive disruption and introduction of restriction sites.

[0242] For disruption of restriction sites, Sal I: (sense strand) 5'-ggagaagtctcaacaccgtccaccaagataatcgatcag-3' (SEQ ID NO: 35), (antisense strand) 5'-ctgatcgattatcttgggtggacgggtgttgagacttctcc-3' (SEQ ID NO: 36), Nhe I: (sense strand) 5'-gtatagtgttcagttgagctgtctcggtctaaggc-3' (SEQ ID NO: 37), (antisense strand) 5'-gccttagaccgacagcaagctcaactgaacacataac-3' (SEQ ID NO: 38), Xho I: (sense strand) 5'-caatgaactctctagagaggctggagtcactaaagagttacctqg-3' (SEQ ID NO: 39), (antisense strand) 5'-ccaggtaacctcttagtgactccagcctctctagagagttcattg-3' (SEQ ID NO: 40), and for introducing restriction sites, NP-P: (sense strand) 5'-gtgaaagttcaccacgatcggtcactcgaggccacaccccaaccacg-3' (SEQ ID NO: 41), (antisense strand) 5'-cgggtgggtgtgggtggtggtgagtgagccgatcggtggatgaactttcac-3' (SEQ ID NO: 42), P-M: (sense strand) 5'-cttaggggtgaaagaaatttcagctagcagcgcgcaatggcagatc-3' (SEQ ID NO: 43), (antisense strand) 5'-gatatctgcatcgccgtgctagctgaaatttcttccaccctaaag-3' (SEQ ID NO: 44), M-F: (sense strand) 5'-cttaggggataaagtcctgtgcgcgtgtgttgcaaaactctcccc-3' (SEQ ID NO: 45), (antisense strand) 5'-ggggagagttttgcaaccaagcgcgacaaagggactttatcctaaag-3' (SEQ ID NO: 46), F-HN: (sense strand) 5'-ggtcgcgcgttacttttagtcgacacctaacaagcacagatcatgg-3' (SEQ ID NO: 47), (antisense strand) 5'-ccatgatctgtgctgtgttgagggtgctgactaaagtaccgcgcgacc-3' (SEQ ID NO: 48), HN-L: (sense strand) 5'-cccagggtgaatgggaaggccggccagggtcatggatggcaggagttcc-3' (SEQ ID NO: 49), (antisense strand) 5'-ggactctgccccatcatgacctggccgcccctccattcaccctggg-3' (SEQ ID NO: 50), were synthesized and used for the reaction. After introduction, each fragment was recovered and purified similarly as described above, and cDNA were assembled.

[0243] In the case of method 2, (sense strand) 5'-ggccgcttaataacgggtttaaaccgcgccaacagtggttgataagaaaaacttaggggtgaagttcatcac-3' (SEQ ID NO: 51), (antisense strand) 5'-ggccgctgatgaactttcaccctaaagttttctatcaaacactgttggcgcggtttaaaccgttaataagc-3' (SEQ ID NO: 52), were synthesized, and after phosphorylation, annealed by 85°C 2 min, 65°C 15 min, 37°C 15 min, and room temperature 15 min to incorporate into SeV cDNA. Alternatively, multicloning sites of pUC18 or pBluescriptII, or the like, are subcloned by PCR using primers comprising termination signal - intervening sequence - initiation signal and then incorporate the resultant into SeV cDNA. The virus reconstitution by resultant cDNA can be performed as described above.

Industrial Applicability

[0244] The present invention provided Paramyxoviridae virus-derived RNP deficient in at least one envelope gene, and the utilization thereof as a vector. As a preferable embodiment, vectors comprising a complex of RNP and a cationic compound are provided. By applying present invention, antigenicity and/or cytotoxicity problems can be avoided when introducing vectors into target cells.

SEQUENCE LISTING

5 <110> Dनावेक Research Inc.

10 <120> PARAMYXOVIRUS-DERIVED RNP

15 <130> D3-102PCT

 <140>

 <141>

20 <150> JP 1999-200740

 <151> 1999-05-18

25 <160> 52.

30 <170> PatentIn Ver. 2.0

35 <210> 1

 <211> 18

 <212> DNA

40 <213> Artificial Sequence

45 <220>

 <223> Description of Artificial Sequence: Artificially
 Synthesized Sequence

50

55

<400> 1

atgcatgccg gcagatga

18

<210> 2

<211> 18

<212> DNA

<213> Artificial Sequence

<220>

<223> Description of Artificial Sequence: Artificially
Synthesized Primer Sequence

<400> 2

gttgagtact gcagagc

18

<210> 3

<211> 42

<212> DNA

<213> Artificial Sequence

<220>

<223> Description of Artificial Sequence: Artificially
Synthesized Primer Sequence

<400> 3

lllgccggca tgcattttc ocaagggag agttllgcaa cc

42

5

<210> 4

<211> 18

<212> DNA

10

<213> Artificial Sequence

15

<220>

<223> Description of Artificial Sequence: Artificially

Synthesized Primer Sequence

20

<400> 4

atgcatgctg gcagatga

25

18

<210> 5

<211> 21

<212> DNA

<213> Artificial Sequence

35

<220>

<223> Description of Artificial Sequence: Artificially

Synthesized Primer Sequence

40

<400> 5

tgggtgaatg agagaatcag c

45

21

50

55

<210> 6

<211> 30

<212> DNA

<213> Artificial Sequence

<220>

<223> Description of Artificial Sequence: Artificially

Synthesized Primer Sequence

<400> 6

atgcatatgg tgatgcggtt ttggcagtac

30

<210> 7

<211> 30

<212> DNA

<213> Artificial Sequence

<220>

<223> Description of Artificial Sequence: Artificially

Synthesized Primer Sequence

<400> 7

tgccggctat tattacttgt acagetcgtc

30

<210> 8

<211> 21

<212> DNA

<213> Artificial Sequence

5

<220>

<223> Description of Artificial Sequence: Artificially
Synthesized Primer Sequence

10

<400> 8

atcagagacc tgcgacaatg c

21

15

20

<210> 9

<211> 21

<212> DNA

25

<213> Artificial Sequence

<220>

<223> Description of Artificial Sequence: Artificially
Synthesized Primer Sequence

30

35

<400> 9

aagtcgtgct gcttcattgt g

21

40

<210> 10

<211> 25

<212> DNA

<213> Artificial Sequence

45

50

55

<220>

5

<223> Description of Artificial Sequence: Artificially
Synthesized Primer Sequence

10

<400> 10

acaaccacta cctgagcacc cagtc

25

15

<210> 11

<211> 21

20

<212> DNA

<213> Artificial Sequence

25

<220>

<223> Description of Artificial Sequence: Artificially
Synthesized Primer Sequence

30

<400> 11

35

gcctaacaca tccagagatc g

21

<210> 12

40

<211> 20

<212> DNA

45

<213> Artificial Sequence

<220>

50

55

<223> Description of Artificial Sequence: Artificially
Synthesized Primer Sequence

5

<400> 12

10

acattcatga gtcagcclgc

20

<210> 13

15

<211> 21

<212> DNA

<213> Artificial Sequence

20

<220>

<223> Description of Artificial Sequence: Artificially
Synthesized Primer Sequence

25

<400> 13

30

atcagagacc igcgacaalg c

21

35

<210> 14

<211> 21

<212> DNA

40

<213> Artificial Sequence

45

<220>

<223> Description of Artificial Sequence: Artificially
Synthesized Primer Sequence

50

55

<400> 14

5

uagtcgtgct gcttcaatg g

21

<210> 15

10

<211> 23

<212> DNA

15

<213> Artificial Sequence

<220>

20

<223> Description of Artificial Sequence: Artificially
Synthesized Primer Sequence

25

<400> 15

gaataaccta gggataaagt ccc

23

30

<210> 16

<211> 19

35

<212> DNA

<213> Artificial Sequence

40

<220>

<223> Description of Artificial Sequence: Artificially
Synthesized Primer Sequence

45

<400> 16

50

55

ggtatctcgc ggtatgtgc

19

5

<210> 17

<211> 45

<212> DNA

10

<213> Artificial Sequence

15

<220>

<223> Description of Artificial Sequence: artificially

synthesized sequence

20

<400> 17

gcggggcgc cgtacgggtg caaccatgc gttactttg accaa

45

25

<210> 18

<211> 80

30

<212> DNA

<213> Artificial Sequence

35

<220>

<223> Description of Artificial Sequence: artificially

synthesized sequence

40

<400> 18

gcggggcgc gctgaacttt caccctaagc tttcttact acggcgtagc ctattacttc 60

tgacaacaga caactggta

80

50

55

<210> 19

5

<211> 41

<212> DNA

<213> Artificial Sequence

10

<220>

15

<223> Description of Artificial Sequence: artificially
synthesized sequence

20

<400> 19

ccaccgacca caccagcgg ccgcgacagc cacggcttcg g

41

25

<210> 20

<211> 41

30

<212> DNA

<213> Artificial Sequence

35

<220>

<223> Description of Artificial Sequence: artificially
synthesized sequence

40

<400> 20

45

ccgagccgt ggcgcgcgcg gcgcgcgcgc ggcgcgcgcg g

41

<210> 21

50

55

<211> 40

<212> DNA

<213> Artificial Sequence

<220>

<223> Description of Artificial Sequence: artificially
synthesized sequence

<400> 21

gaaatttcac ctaagcggcc gcaatggcag atatctatag

40 ..

<210> 22

<211> 40

<212> DNA

<213> Artificial Sequence

<220>

<223> Description of Artificial Sequence: artificially
synthesized sequence

<400> 22

ctatagatat ctgccattgc ggccgcttag gtgaaatttc

40

<210> 23

<211> 43

<212> DNA

<213> Artificial Sequence

5

<220>

<223> Description of Artificial Sequence: artificially

10

synthesized sequence

<400> 23

15

gggataaagt cccttgcggc cgcttggttg caaaactctc ccc

43

<210> 24

20

<211> 43

<212> DNA

25

<213> Artificial Sequence

<220>

30

<223> Description of Artificial Sequence: artificially

synthesized sequence

35

<400> 24

ggggagagti ttgcaaccaa gggccgcaa gggactttat ccc

43

40

<210> 25

<211> 47

45

<212> DNA

<213> Artificial Sequence

50

55

<220>

<223> Description of Artificial Sequence: artificially
synthesized sequence

<400> 25

ggtcgogogg taotttagcg gccgcctcaa acaagcacag atcatgg

47

<210> 26

<211> 47

<212> DNA

<213> Artificial Sequence

<220>

<223> Description of Artificial Sequence: artificially
synthesized sequence

<400> 26

ccatgatctg tgottgtttg aggcggccgc taaagtaccg cgcgacc

47

<210> 27

<211> 44

<212> DNA

<213> Artificial Sequence

<220>

<223> Description of Artificial Sequence: artificially

synthesized sequence

5 <400> 27
cctgcccatc catgacctag cggccgcttc ccallcaccu tggg 44

10 <210> 28
 <211> 44

15 <212> DNA
 <213> Artificial Sequence

20 <220>
 <223> Description of Artificial Sequence: artificially
 synthesized sequence

25 <400> 28
30 cccagsgtga atgggaagcg gccgctaggt calggatggg cagg 44

 <210> 29
35 <211> 40
 <212> DNA
 <213> Artificial Sequence

40 <220>
 <223> Description of Artificial Sequence: artificially
45 synthesized sequence

50

55

<400> 29

gcggcgcgcc atgctgctgc tgctgctgct gctgggcctg

40

<210> 30

<211> 40

<212> DNA

<213> Artificial Sequence

<220>

<223> Description of Artificial Sequence: artificially
synthesized sequence

<400> 30

gcggcgcgcc cttatcatgt ctgctcgaag cggccggccg

40

<210> 31

<211> 74

<212> DNA

<213> Artificial Sequence

<220>

<223> Description of Artificial Sequence: artificially
synthesized sequence

<400> 31

gcggccgcgt ttaaaccggc gccatttaa atccgtagta agaaaaactt agggtgaaag 60

tccatcgogg ccgc

74

5

<210> 32

<211> 74

10

<212> DNA

<213> Artificial Sequence

15

<220>

<223> Description of Artificial Sequence: artificially

synthesized sequence

20

<400> 32

25

ggcgccgcga tgaactttca ccctaagttt ttcctacac ggatttaast ggcgcgccgt 60

ttaaacgcgg ccgc

74

30

<210> 33

<211> 48

<212> DNA

35

<213> Artificial Sequence

40

<220>

<223> Description of Artificial Sequence: artificially

synthesized sequence

45

<400> 33

acttgcggcc gccaaagttc agtaatgtcc atgttgttct acactctg

48

50

55

<210> 34

5

<211> 72

<212> DNA

<213> Artificial Sequence

10

<220>

<223> Description of Artificial Sequence: artificially
synthesized sequence

15

<400> 34

atccgaggcc gcgatgac taccacclaa gttttctta ctacggtcag cctctcttg 60
tagccttct gc 72

20

25

<210> 35

<211> 40

<212> DNA

<213> Artificial Sequence

30

35

<220>

<223> Description of Artificial Sequence: artificially
synthesized sequence

40

<400> 35

ggagaggtct caacaccgc caccacagat aatcgatcag 40

45

50

55

<210> 36

<211> 40

<212> DNA

<213> Artificial Sequence

<220>

<223> Description of Artificial Sequence: artificially
synthesized sequence

<400> 36

ctgatcgatt atcttgggtg gaagggtgtg agacttctcc

40

<210> 37

<211> 38

<212> DNA

<213> Artificial Sequence

<220>

<223> Description of Artificial Sequence: artificially
synthesized sequence

<400> 37

gtatatgtgt tcagttgagc ttgctgtcgg tctaaggc

38

<210> 38

<211> 38

<212> DNA

<213> Artificial Sequence

5

<220>

<223> Description of Artificial Sequence: artificially
synthesized sequence

10

<400> 38

gccttagacc gacagcaagc tcaac1gaac acatatac

38

15

20

<210> 39

<211> 45

<212> DNA

25

<213> Artificial Sequence

<220>

<223> Description of Artificial Sequence: artificially
synthesized sequence

30

35

<400> 39

cautgaactc tctagagagg ctggagtcac laaagagtta cctgg

45

40

<210> 40

<211> 45

<212> DNA

<213> Artificial Sequence

45

50

55

<220>

5 <223> Description of Artificial Sequence: artificially
 synthesized sequence

10 <400> 40

ccaggtaact ctttagtgac tccagcctct ctagagagtt cattg 45

15 <210> 41
 <211> 52

20 <212> DNA

<213> Artificial Sequence

25 <220>

<223> Description of Artificial Sequence: artificially
 synthesized sequence

30

<400> 41

35 gtgaagttc atccaccgat cggctcactc gaggccacac ccaaccccccac cg 52

40 <210> 42
 <211> 52

<212> DNA

<213> Artificial Sequence

45

<220>

50

55

<223> Description of Artificial Sequence: artificially
synthesized sequence

5

<400> 42

cggtgggggtt ggggtgggcc tggagtggc cgatcggtgg atgaactttc ac 62

10

<210> 43

<211> 47

15

<212> DNA

<213> Artificial Sequence

20

<220>

<223> Description of Artificial Sequence: artificially
synthesized sequence

25

<400> 43

cttaggtga aagaatttc agctagcacg gcgcaatggc agatata 47

30

<210> 44

<211> 47

35

<212> DNA

<213> Artificial Sequence

40

<220>

<223> Description of Artificial Sequence: artificially
synthesized sequence

45

50

55

<400> 44

5

gatatctgcc attgcgcctg gctagctgaa atttctttca ccctaag

47

<210> 45

10

<211> 47

<212> DNA

<213> Artificial Sequence

15

<220>

20

<223> Description of Artificial Sequence: artificially
synthesized sequence

25

<400> 45

cttagggata aagtccttg tgcgcgcttg gttgcaaac tctcccc

47

30

<210> 46

<211> 47

<212> DNA

35

<213> Artificial Sequence

40

<220>

<223> Description of Artificial Sequence: artificially
synthesized sequence

45

<400> 46

50

55

ggggagagtt ttgcaaccaa gcgcgcacaa gggactttat ccctaag

47

5

<210> 47

<211> 47

<212> DNA

10

<213> Artificial Sequence

15

<220>

<223> Description of Artificial Sequence: artificially
synthesized sequence

20

<400> 47

ggtcgcgcgg tacttttagtc gacacctcaa acaagcacag atcatgg

25

47

<210> 48

30

<211> 47

<212> DNA

<213> Artificial Sequence

35

<220>

<223> Description of Artificial Sequence: artificially
synthesized sequence

40

<400> 48

ccatgatctg tgcttggttg aggtgtcgac taaagtacog cgcgacc

45

47

50

55

<210> 49

<211> 49

5

<212> DNA

<213> Artificial Sequence

10

<220>

<223> Description of Artificial Sequence: artificially

15

synthesized sequence

<400> 49

20

cccagggtga atgggaaggg ccggccaggl calggalggg caggaglcc

49

<210> 50

25

<211> 49

<212> DNA

30

<213> Artificial Sequence

<220>

35

<223> Description of Artificial Sequence: artificially

synthesized sequence

40

<400> 50

ggaciccgc cccccaatga cctggccggc ccttcccatl caccctggg

49

45

<210> 51

<211> 72

50

55

<212> DNA

<213> Artificial Sequence

<220>

<223> Description of Artificial Sequence: artificially
synthesized sequence

<400> 51

ggccgcttaa ttaacggtt aaacgcgc caacagtgt gataagaaa acttaggg 60
aaagttcatc ac 72

<210> 52

<211> 72

<212> DNA

<213> Artificial Sequence

<220>

<223> Description of Artificial Sequence: artificially
synthesized sequence

<400> 52

ggccgtgatg aactttcacc ctaagtttt cttatcaaca ctgttggcgc gcgtttaaac 60
cgtaattaa gc 72

Claims

1. A complex comprising (a) a negative-strand single-stranded RNA derived from a paramyxovirus, wherein said RNA is modified so as not express at least one of the envelope proteins of Paramyxoviridae viruses, and (b) a

protein encoded by and binding to said negative-strand single-stranded RNA.

2. A complex according to claim 1, wherein said negative-strand single-stranded RNA is modified so as to express NP, P and L proteins, but not F, HN or M proteins, or any combination thereof.
3. A complex according to claim 1 or 2, wherein said negative-strand single-stranded RNA derives from the Sendai virus.
4. A complex according to any of claims 1 through 3, wherein said negative-strand single-stranded RNA further encodes a foreign gene.
5. A composition for gene transfer, comprising a complex according to claim 4 and a cationic lipid.
6. A composition for gene transfer, comprising a complex according to claim 4 and a cationic polymer.
7. A method for expressing a foreign gene in a cell, comprising the step of introducing the composition for gene transfer according to claim 5 or 6 into a cell.

Figure 1

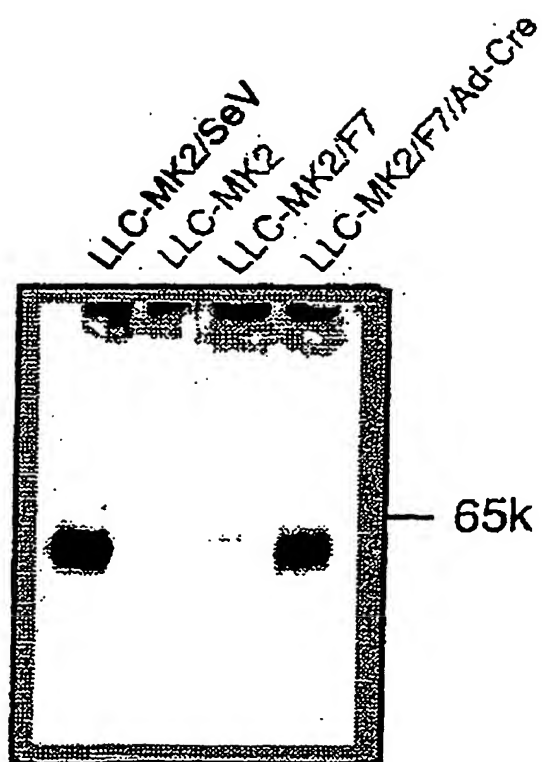


Figure 2

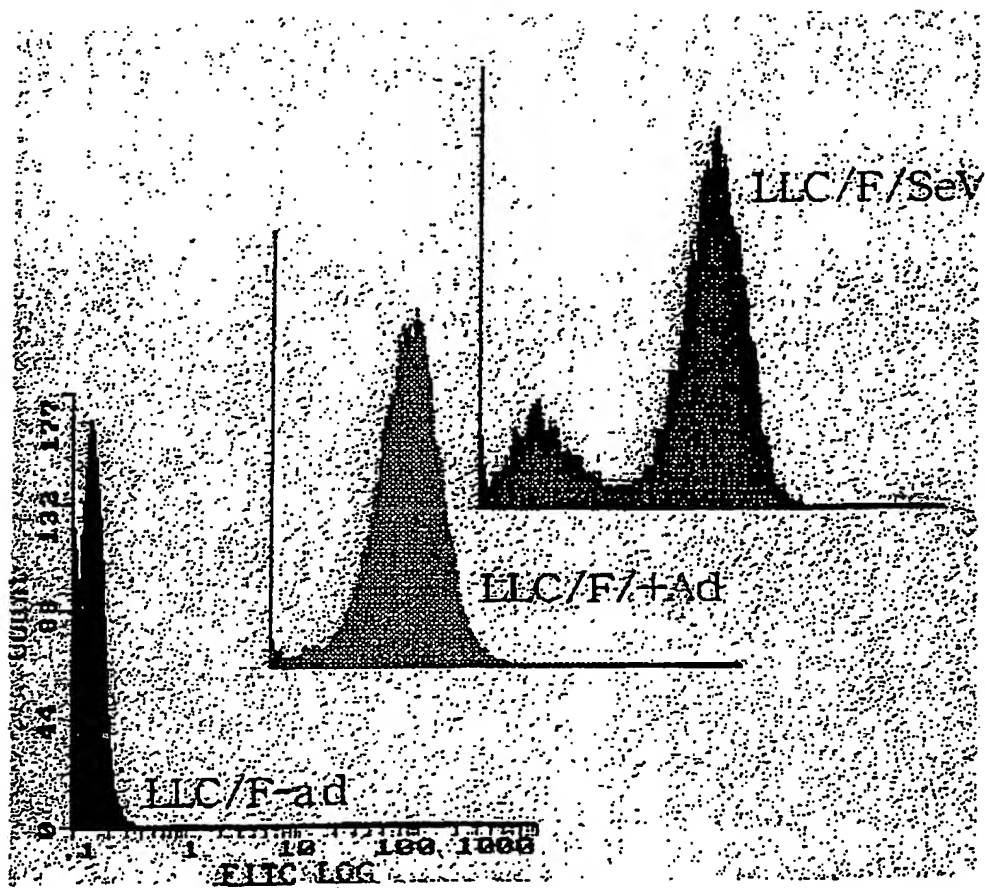


Figure 3

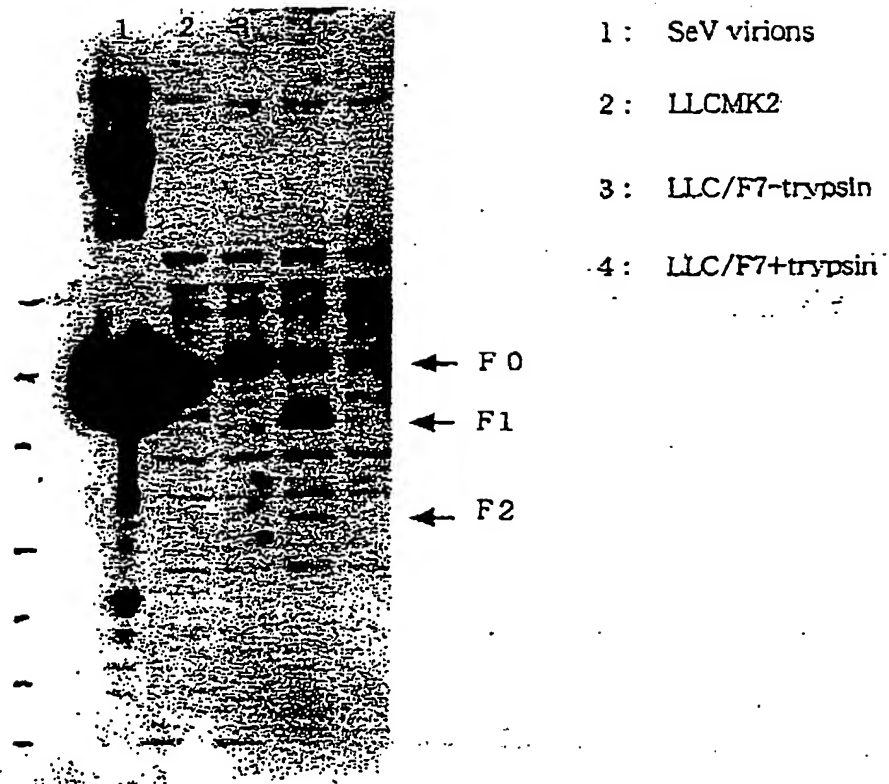


Figure 4

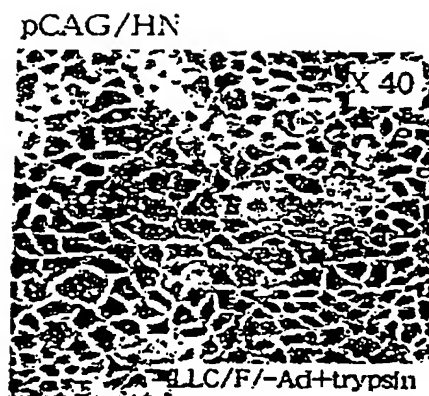
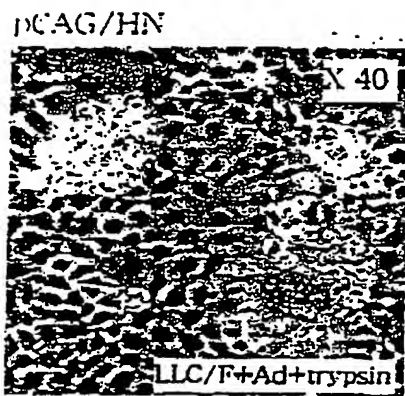


Figure 5

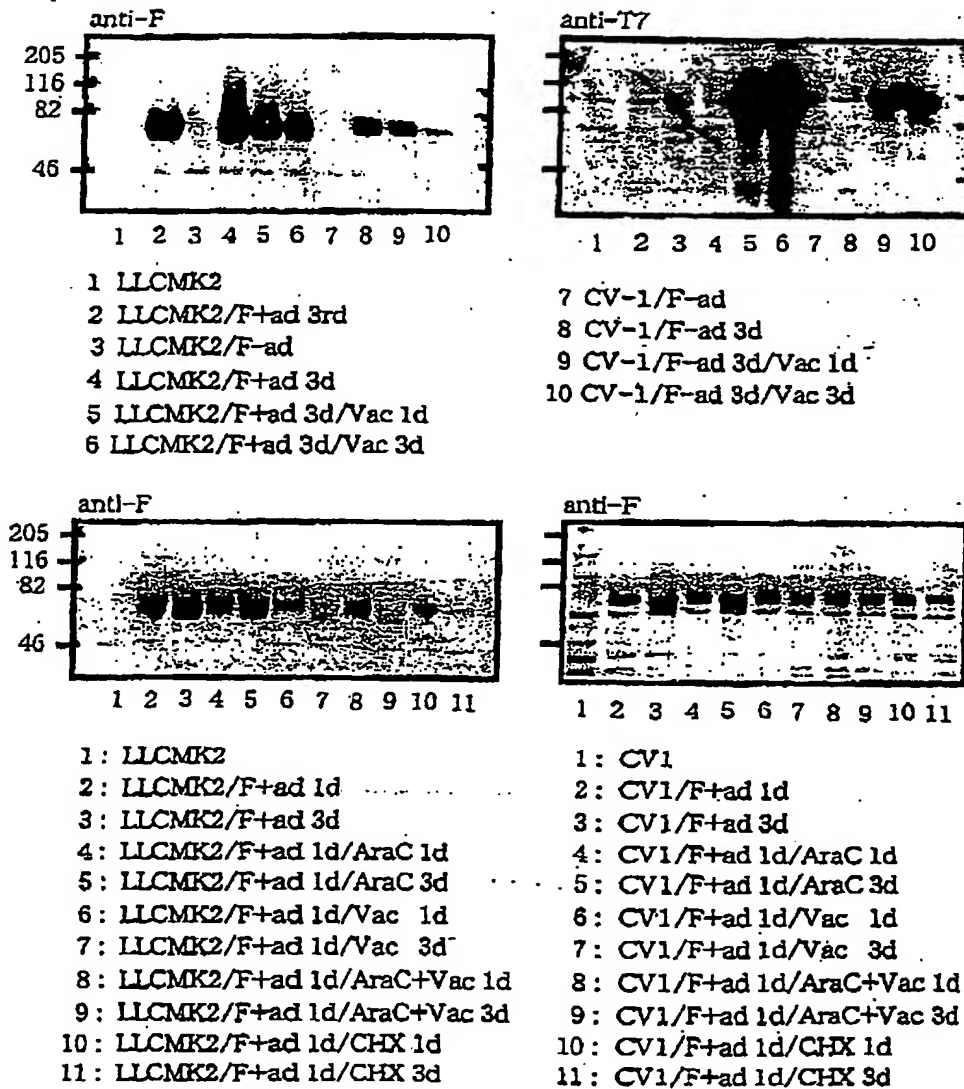


Figure 6

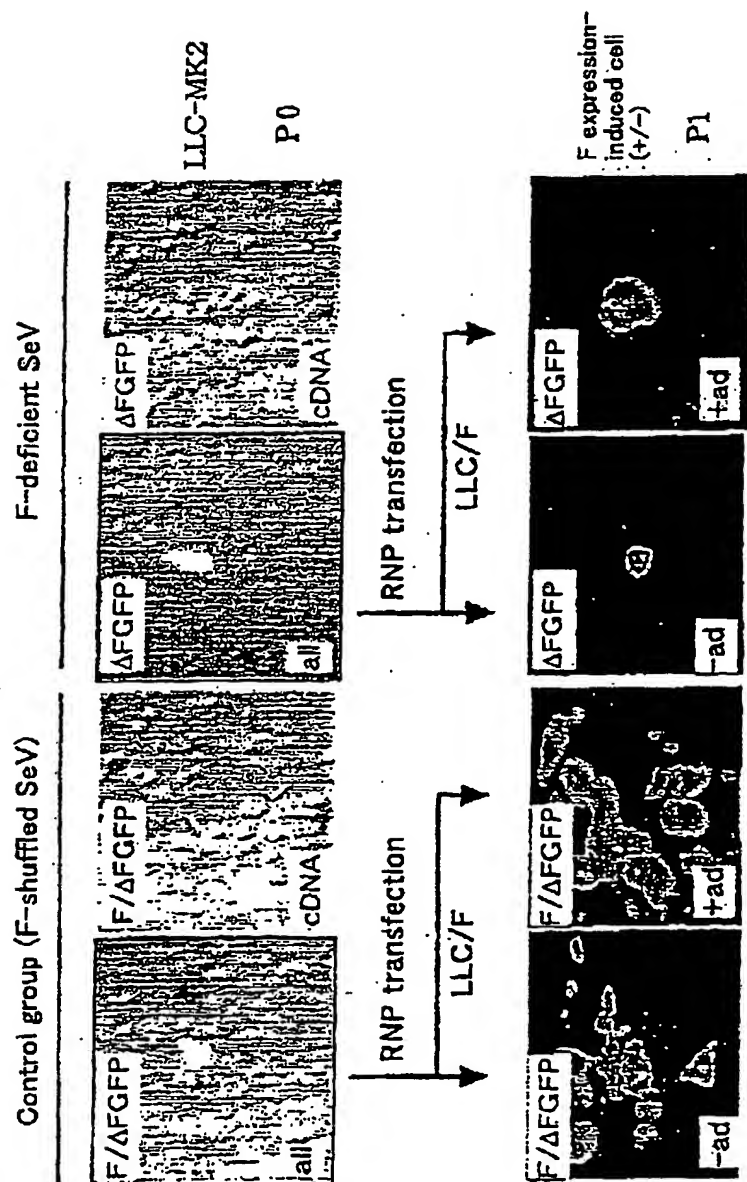


Figure 7

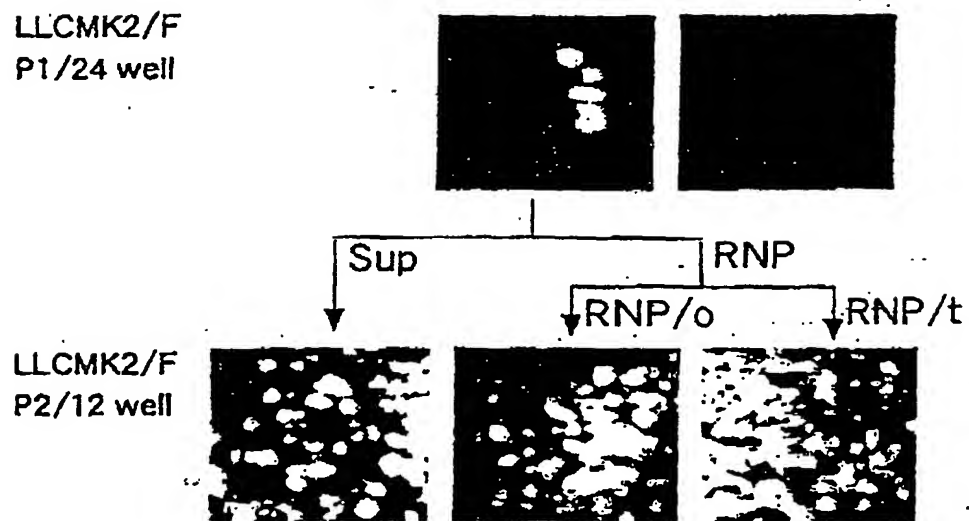


Figure 8

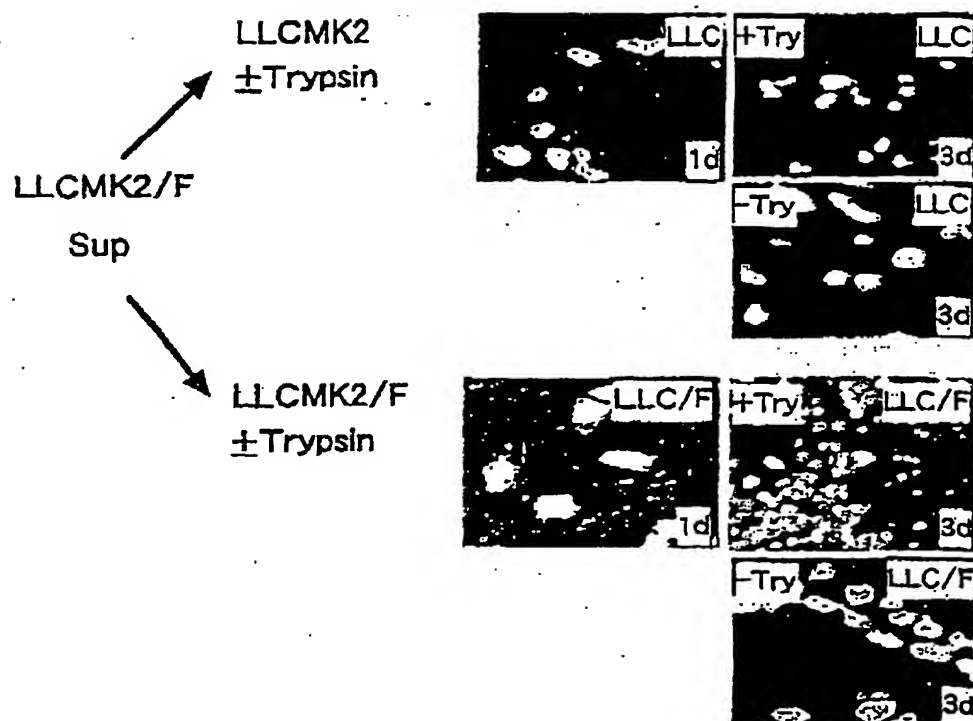
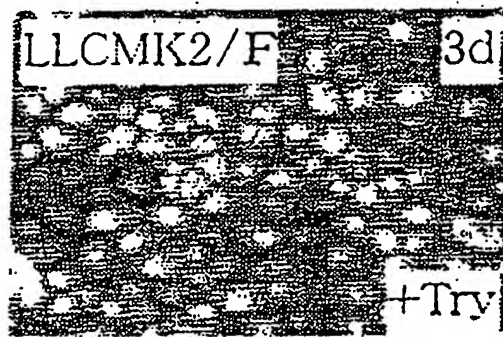


Figure 9

Infection with LLCMK2/F sup



Infection with LLCMK2 sup

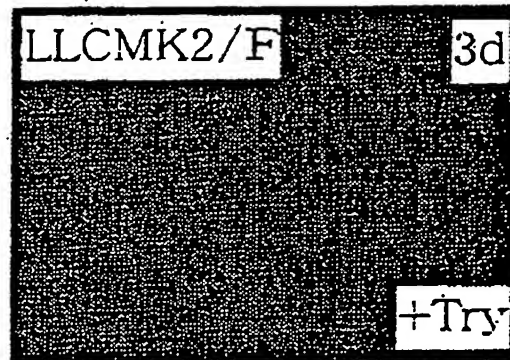


Figure 10

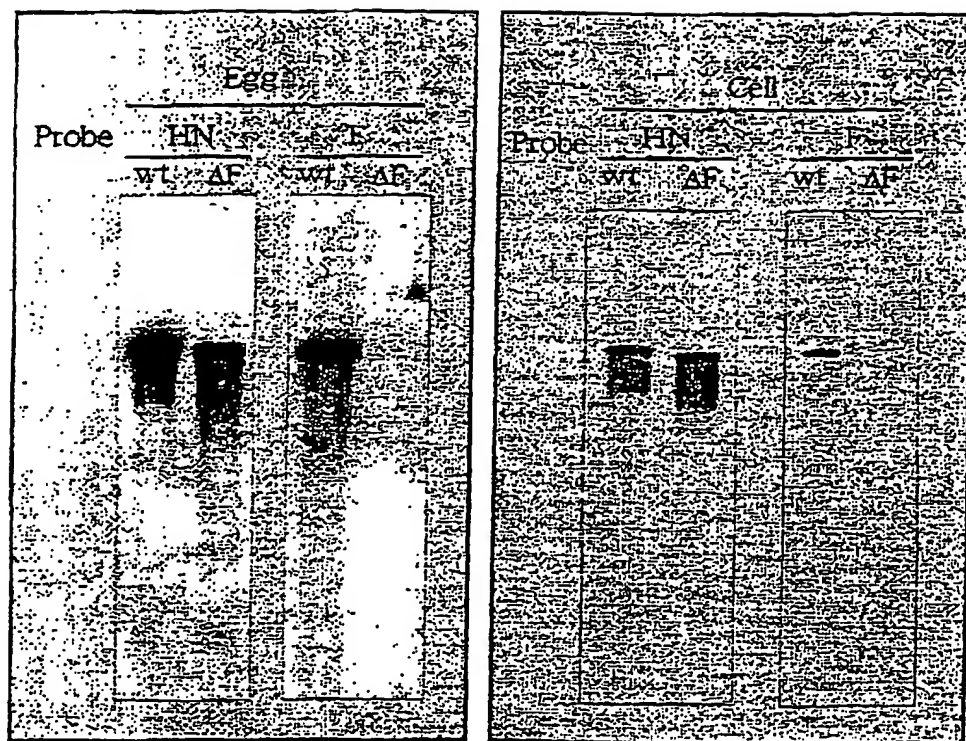
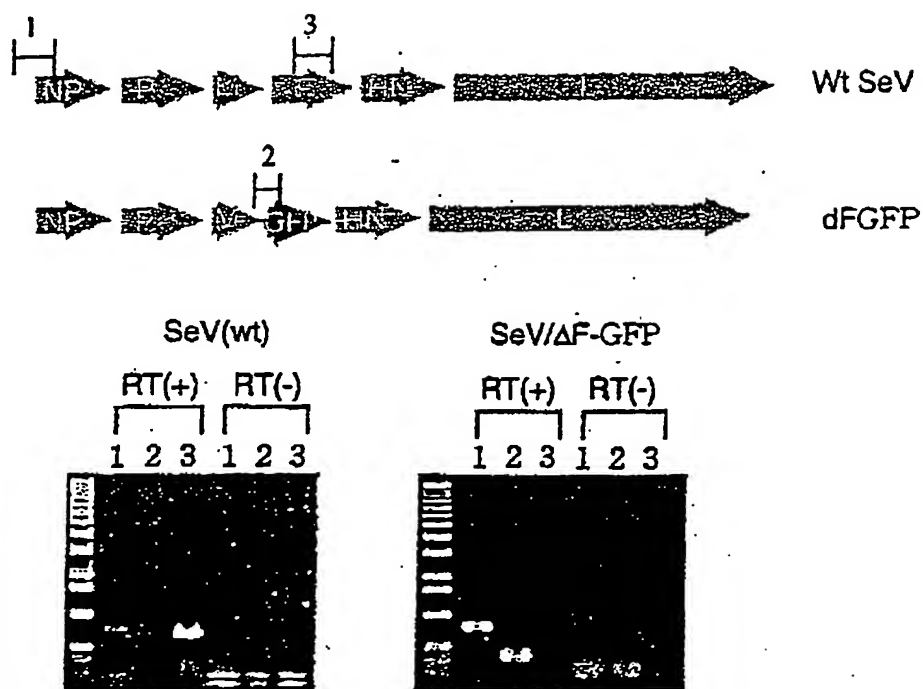


Figure 11



- 1: +18-NP, for the confirmation of the presence of +18 Not I site
- 2: M-GFP, for the confirmation of the presence of the GFP gene in the F gene-deficient region
- 3: F gene, for the confirmation of the presence of the F gene

Figure 12

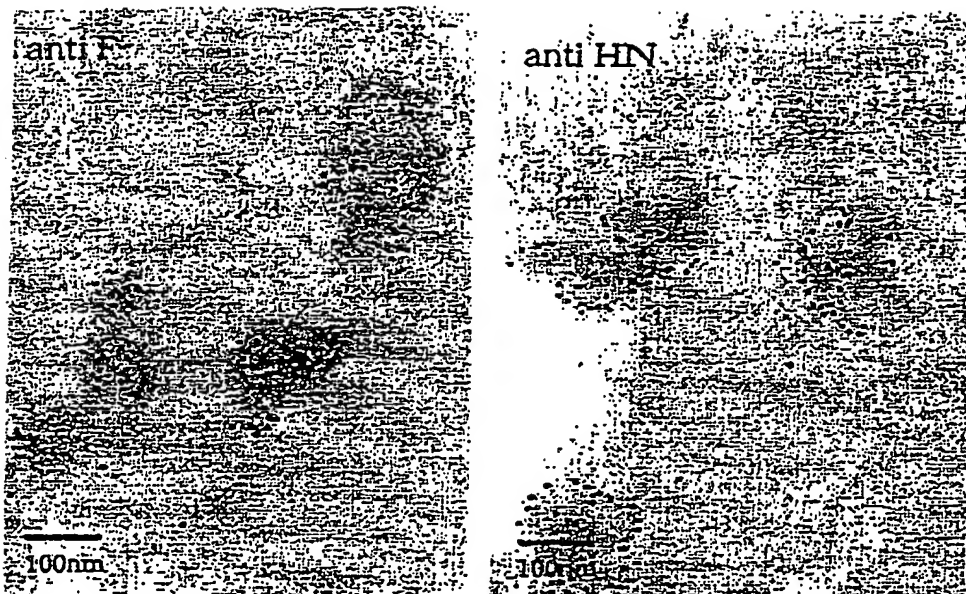
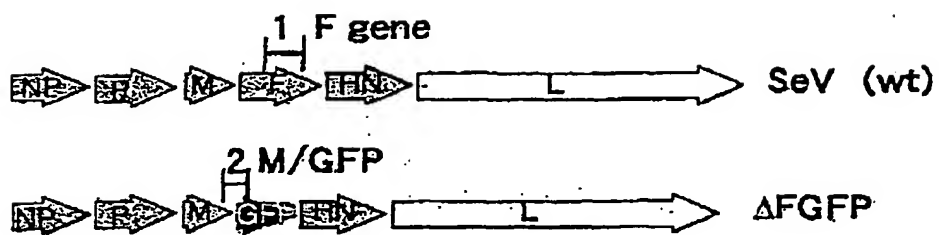


Figure 13



Primer Set	ΔFGFP	F/ΔFGFP	GFP/SeV	SeV (wt)
1	-	+	+	+
2	+	+	-	-

Figure 14

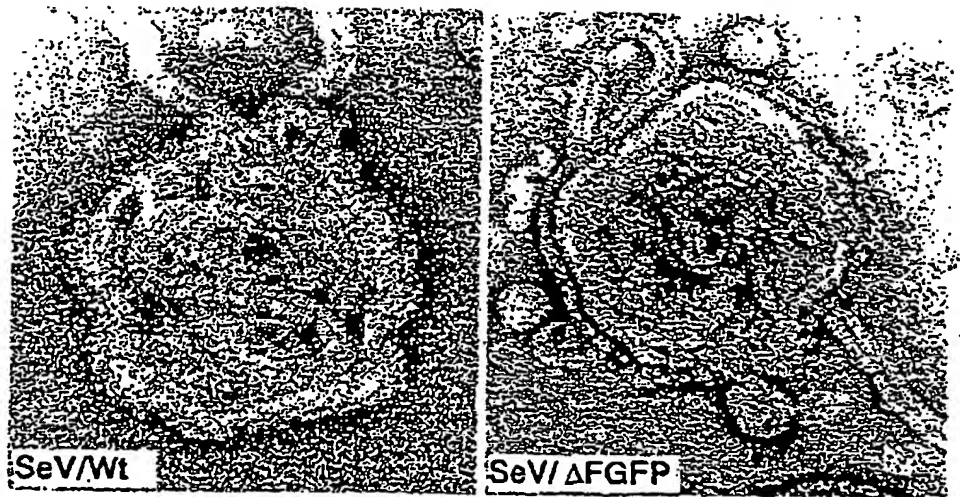


Figure 15

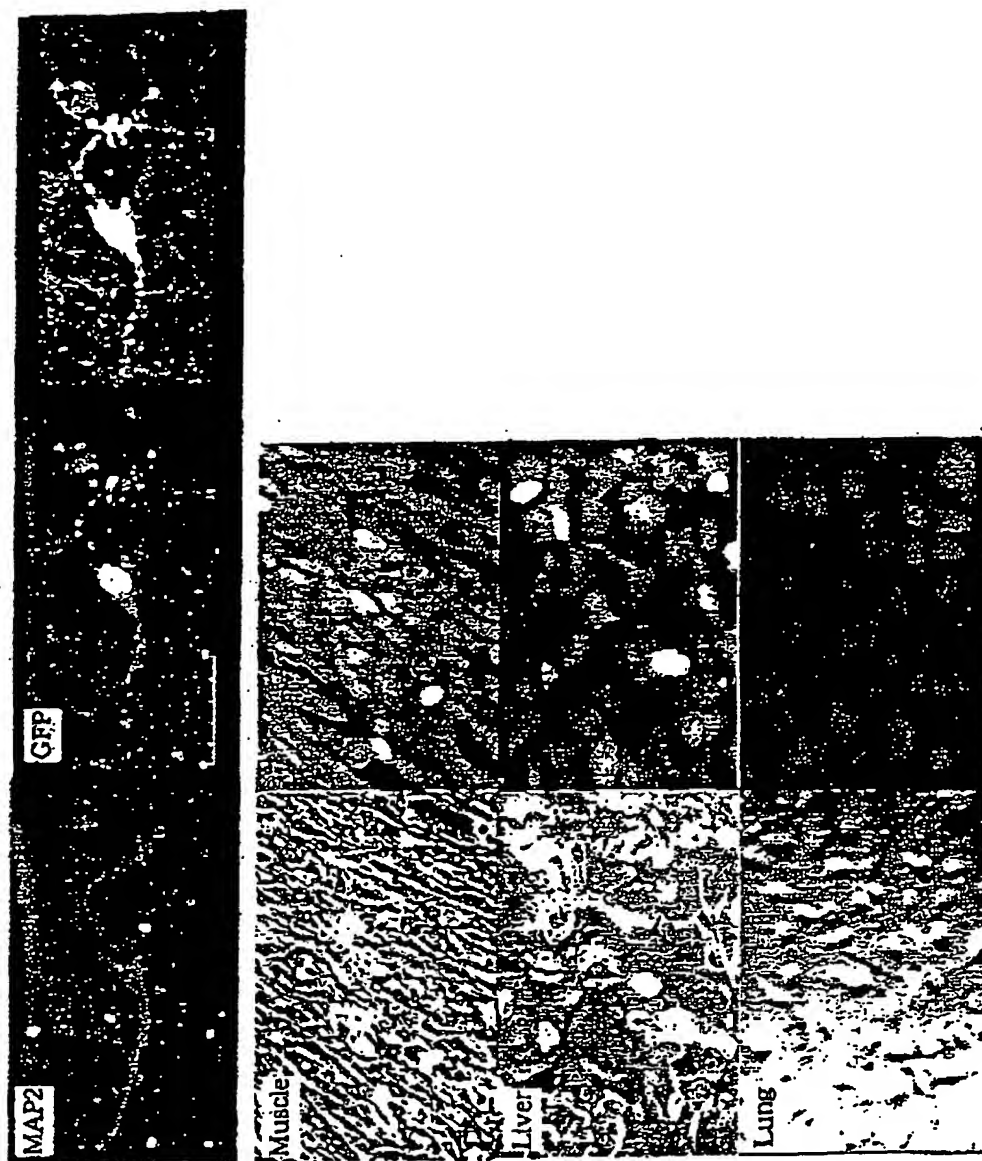


Figure 18

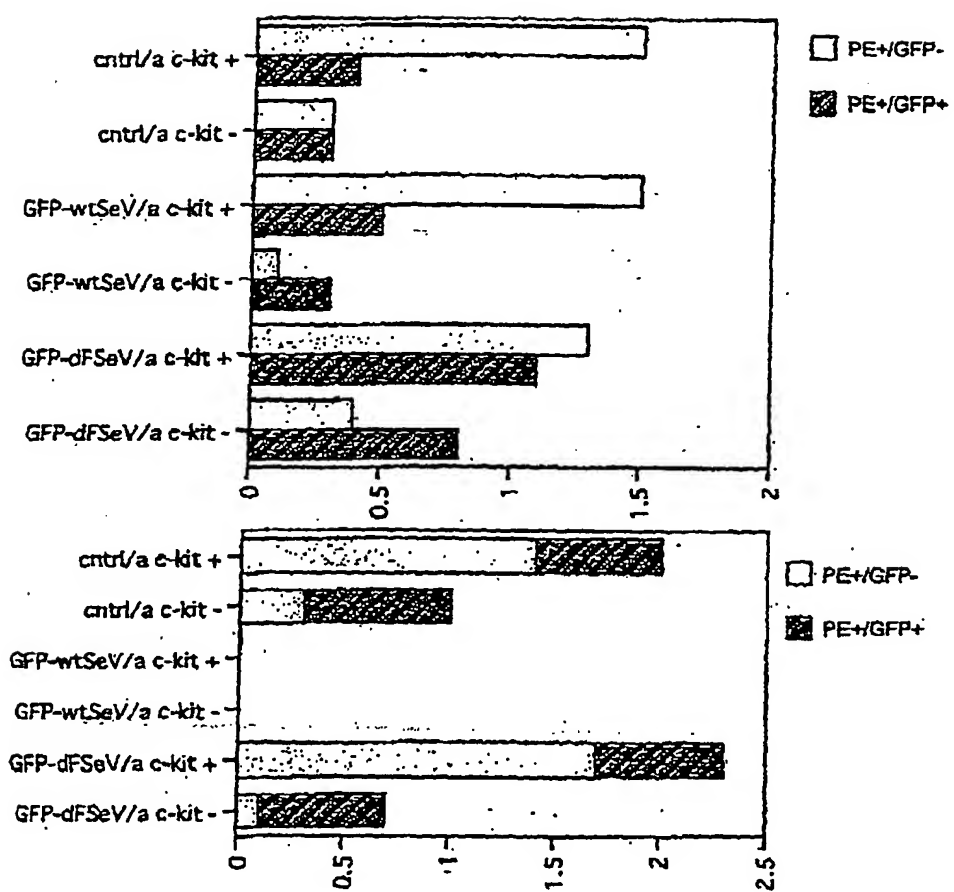


Figure 17

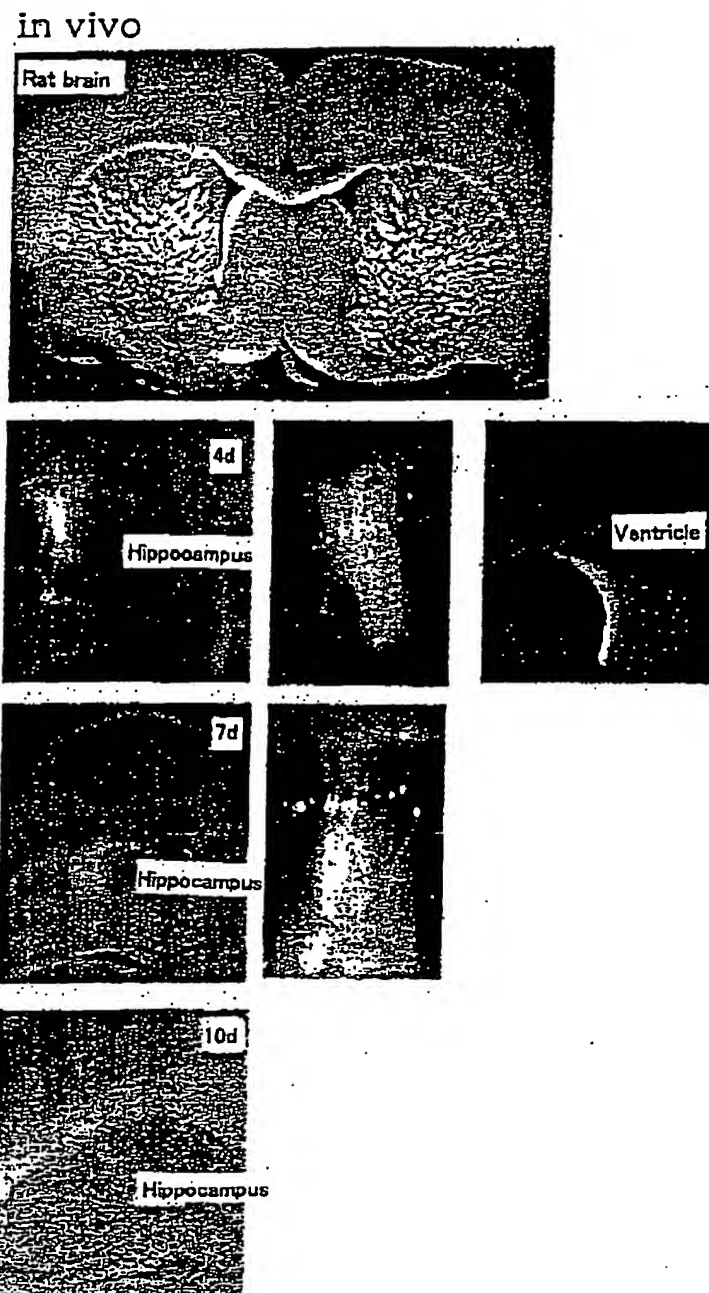


Figure 18

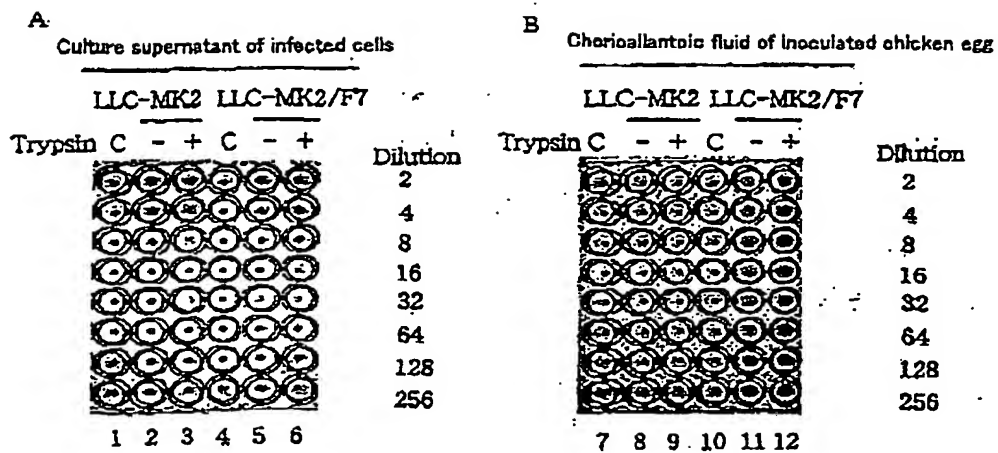


Figure 19

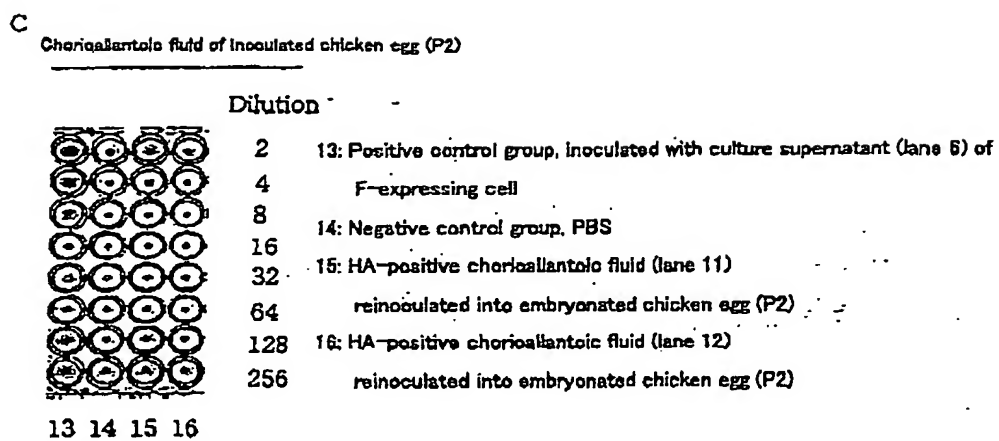


Figure 20

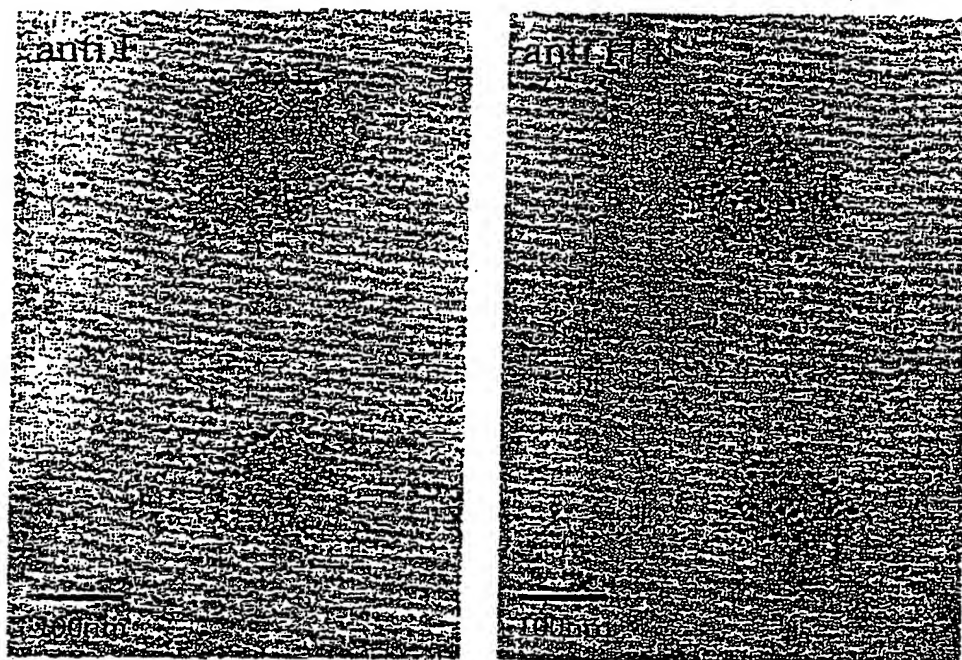


Figure 21

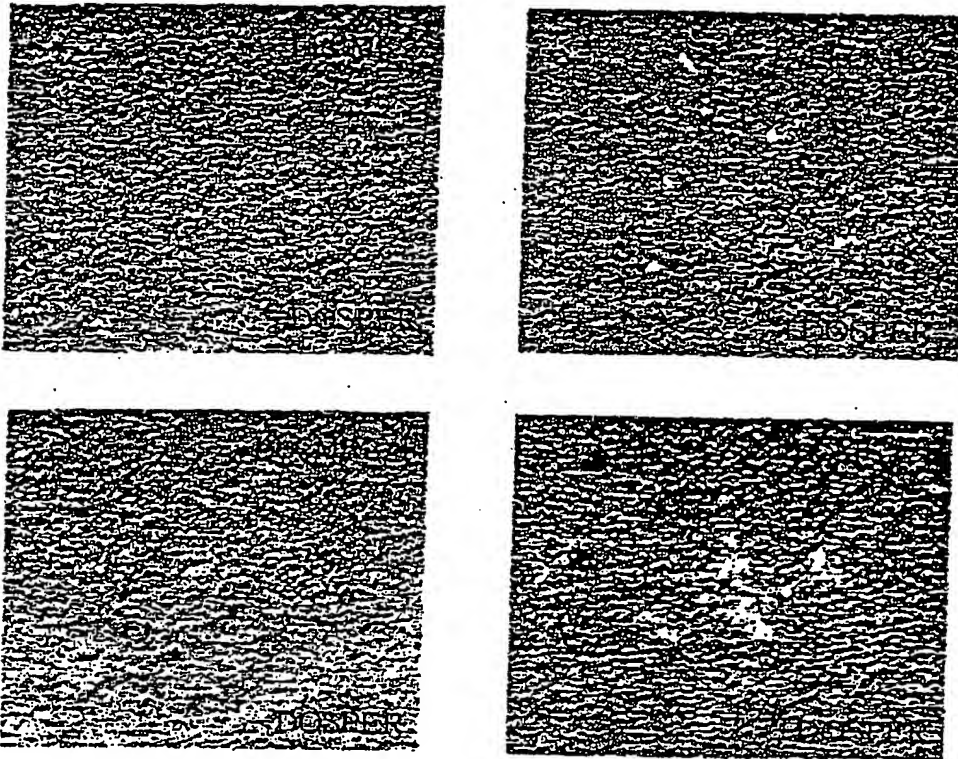


Figure. 22

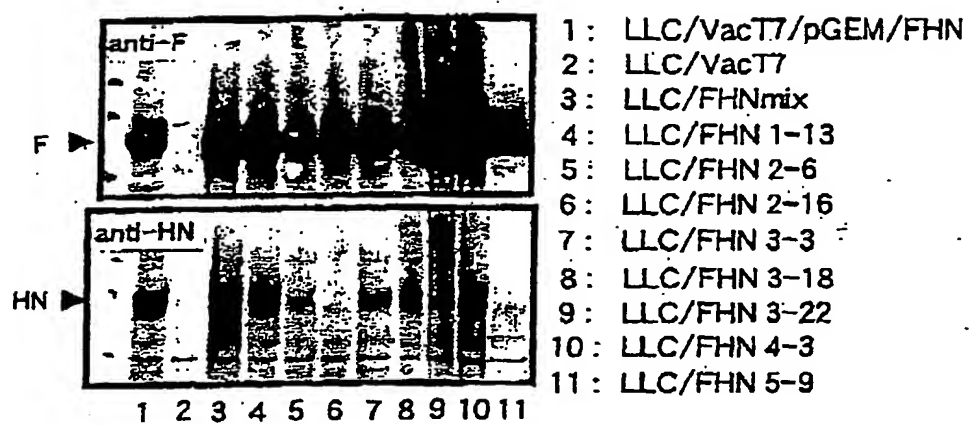


Figure 23

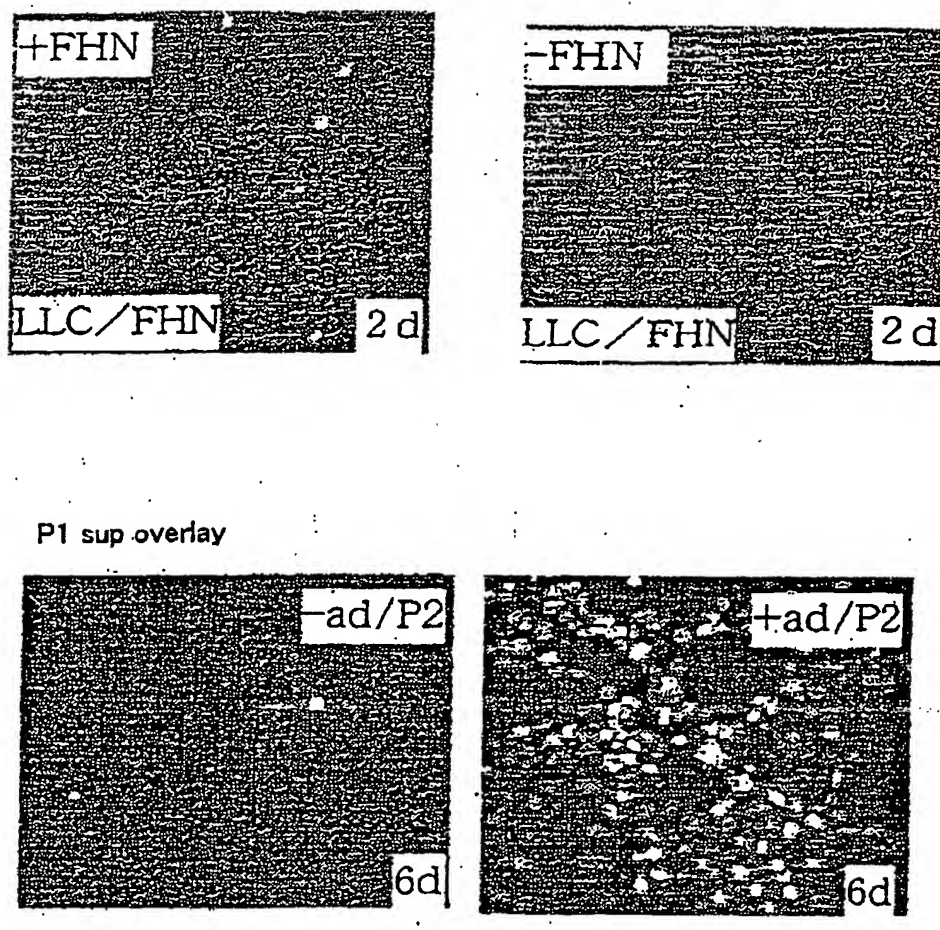


Figure 24

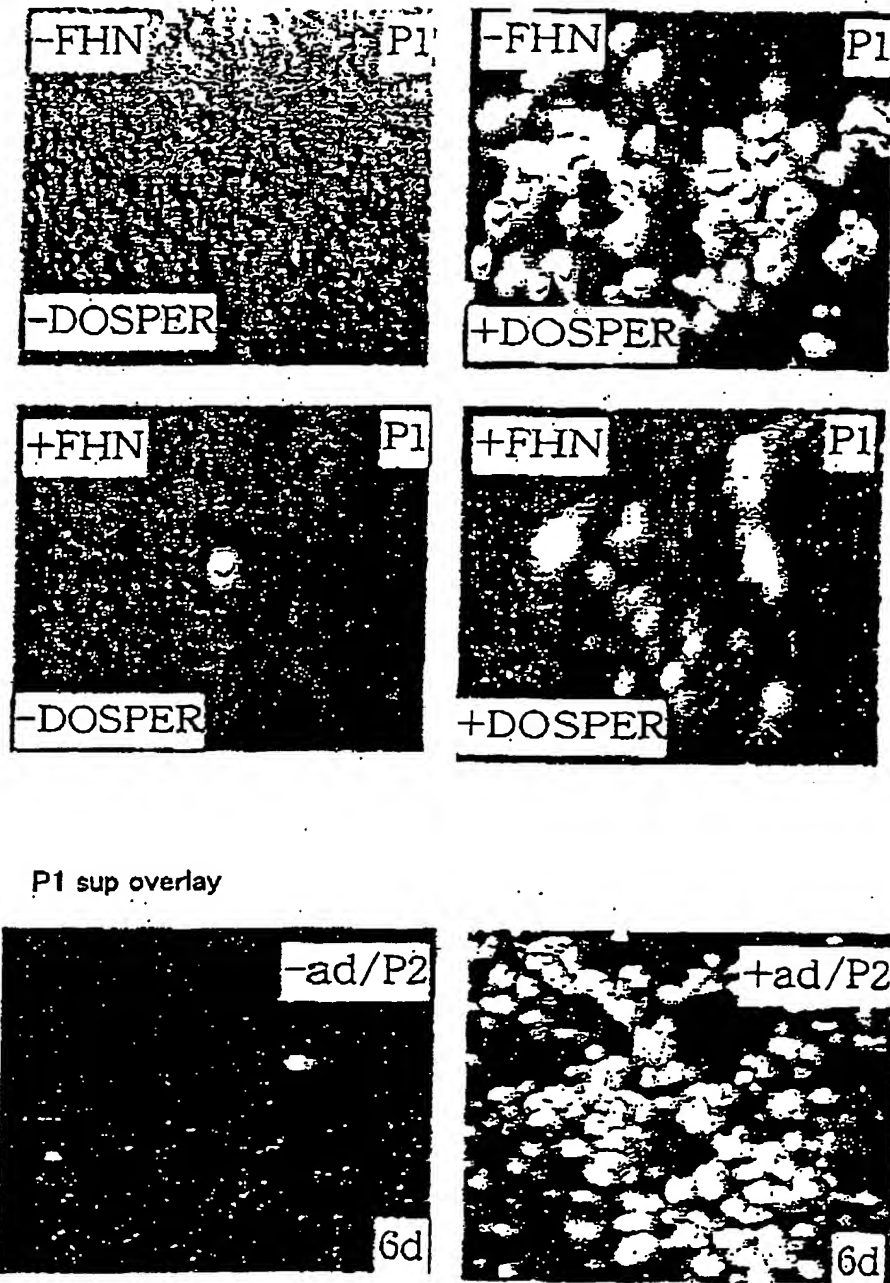


Figure 25

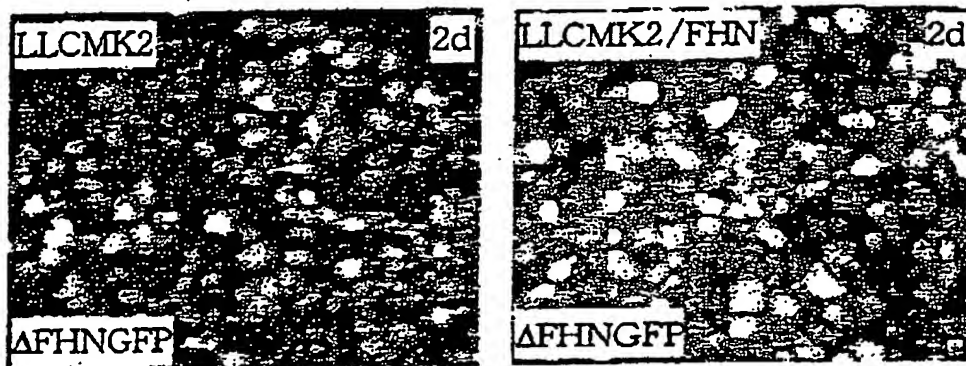
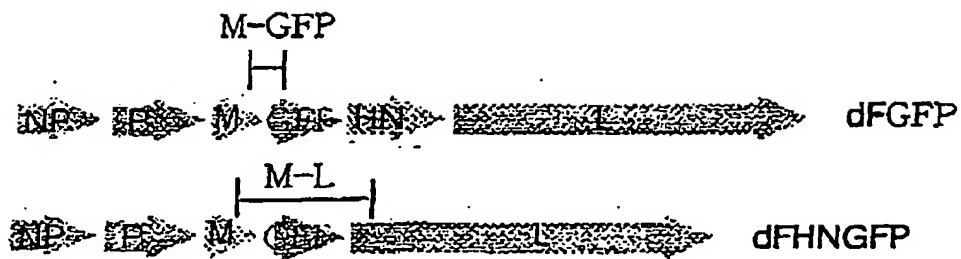


Figure 26



		M-GFP		M-L	
		ΔF		ΔFHN	
		+	-	+	-
RT		+	-	+	-

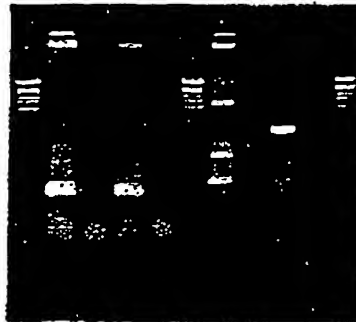


Figure 27

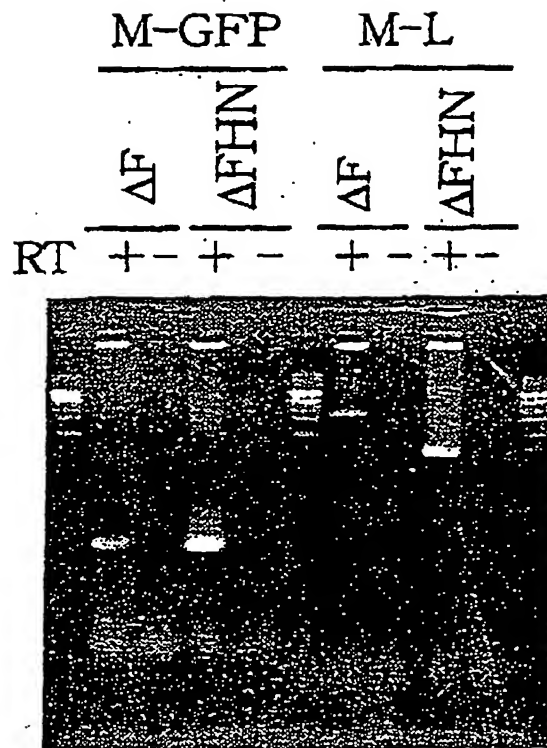
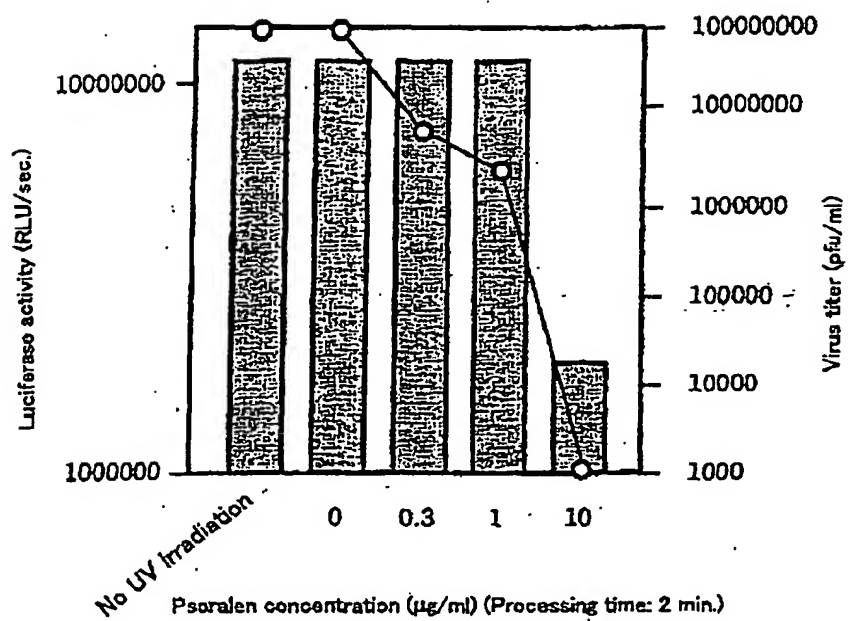


Figure 28



○: Virus titer

Figure 29

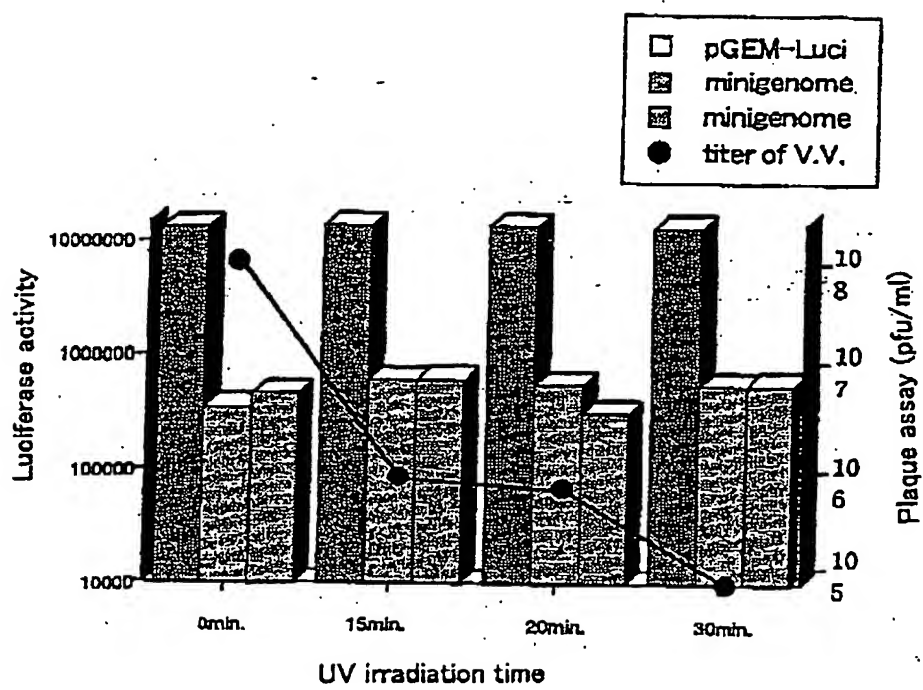


Figure 30

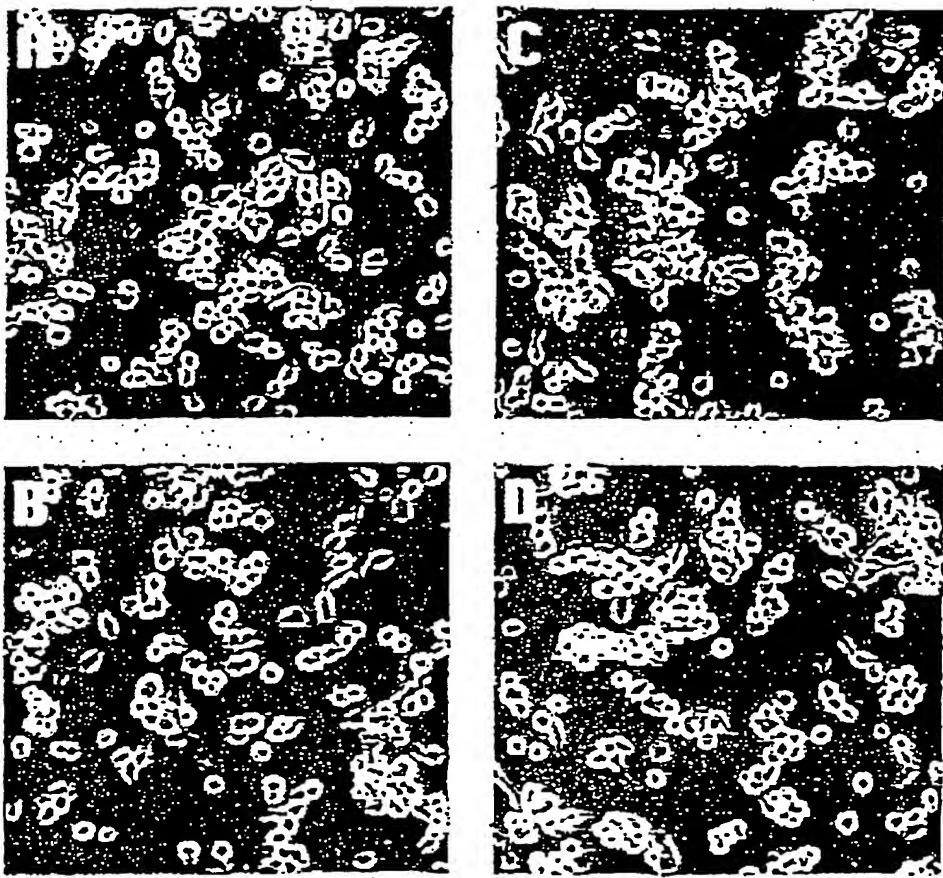


Figure 31

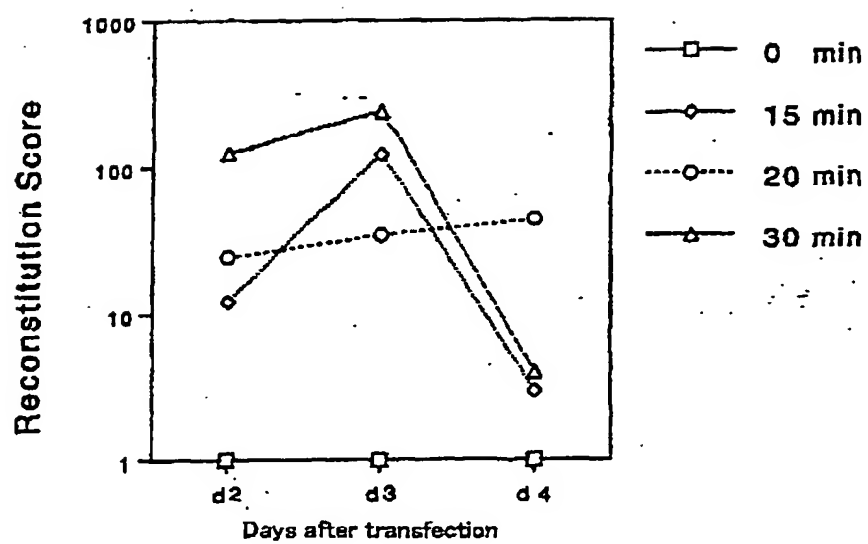


Figure 32

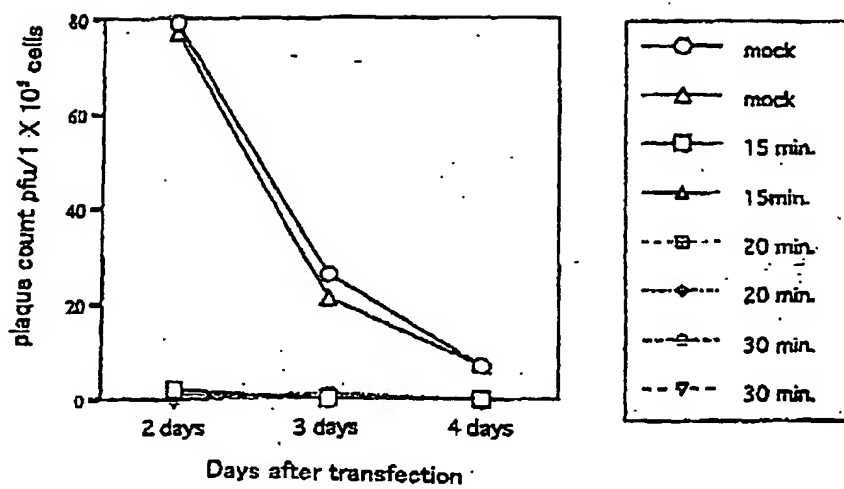


Figure 33

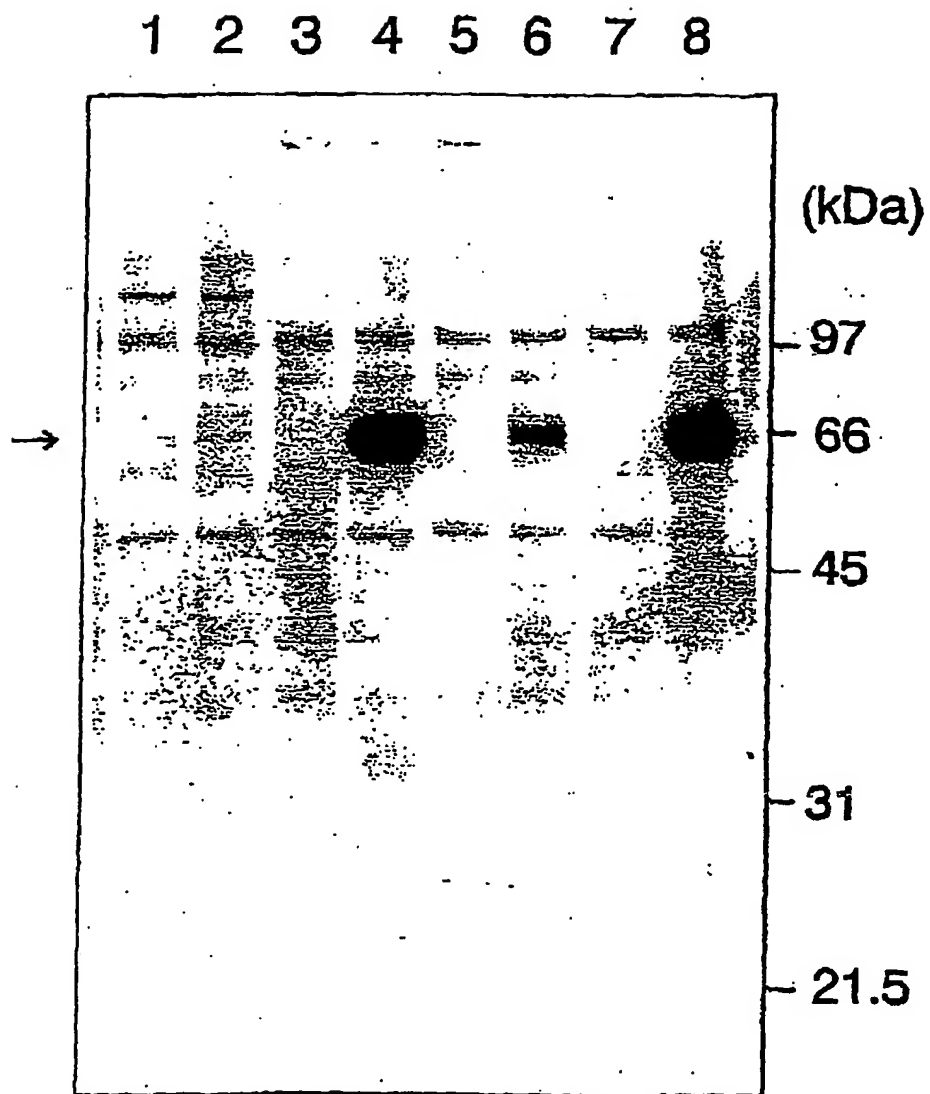


Figure 34

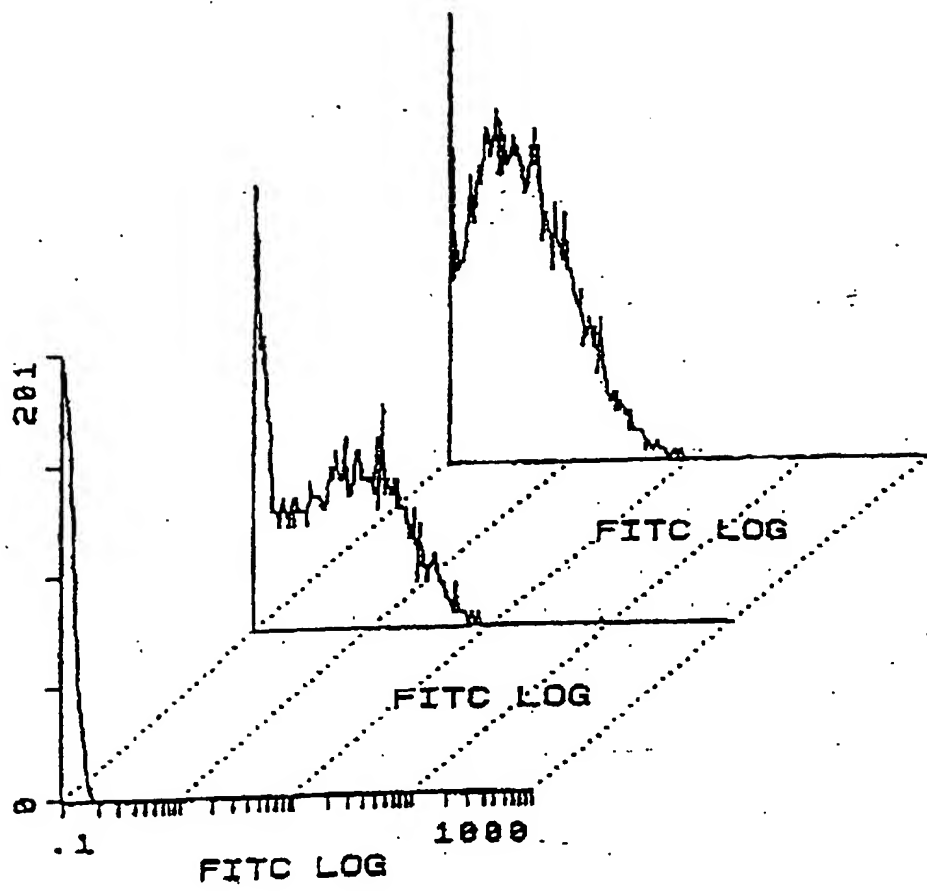


Figure 35

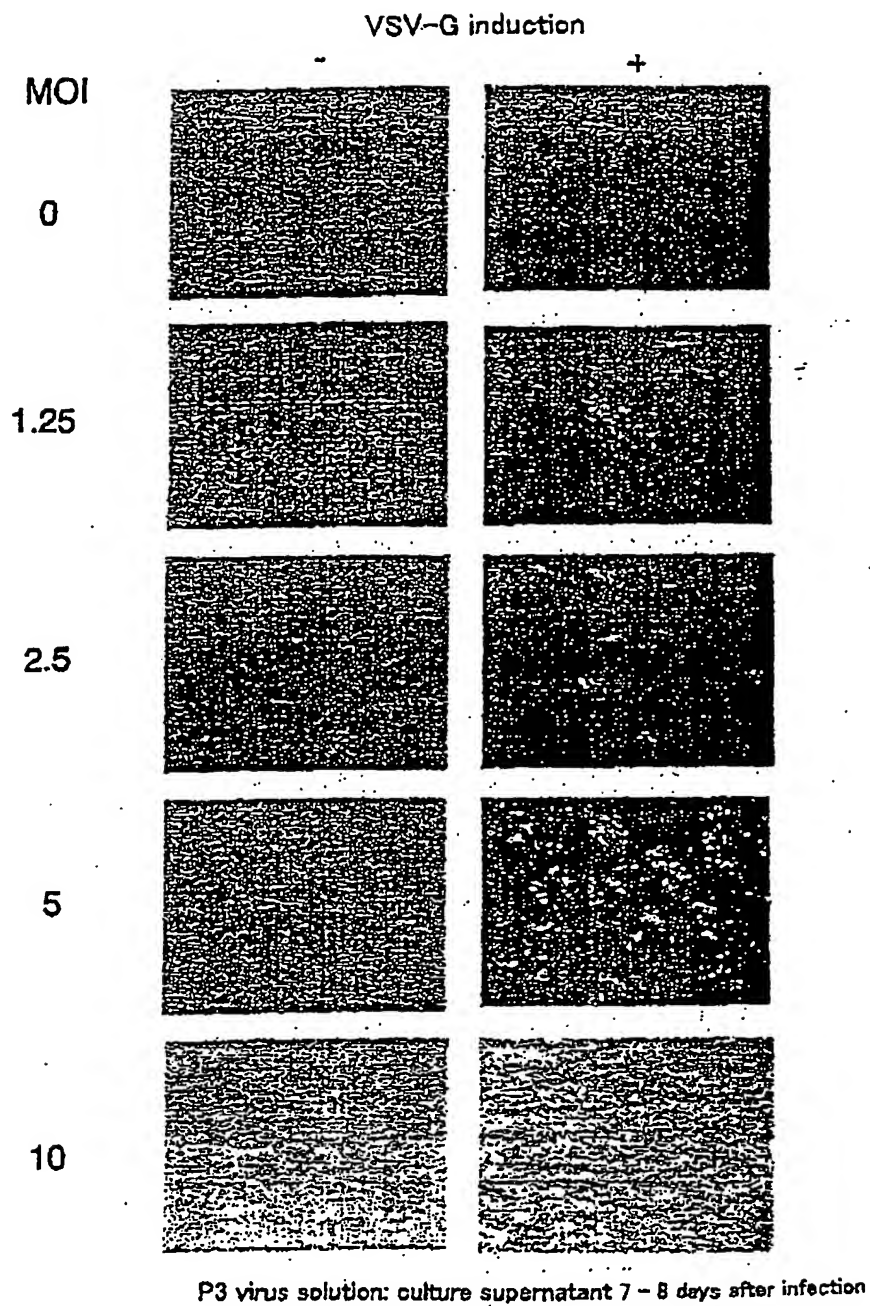


Figure 36

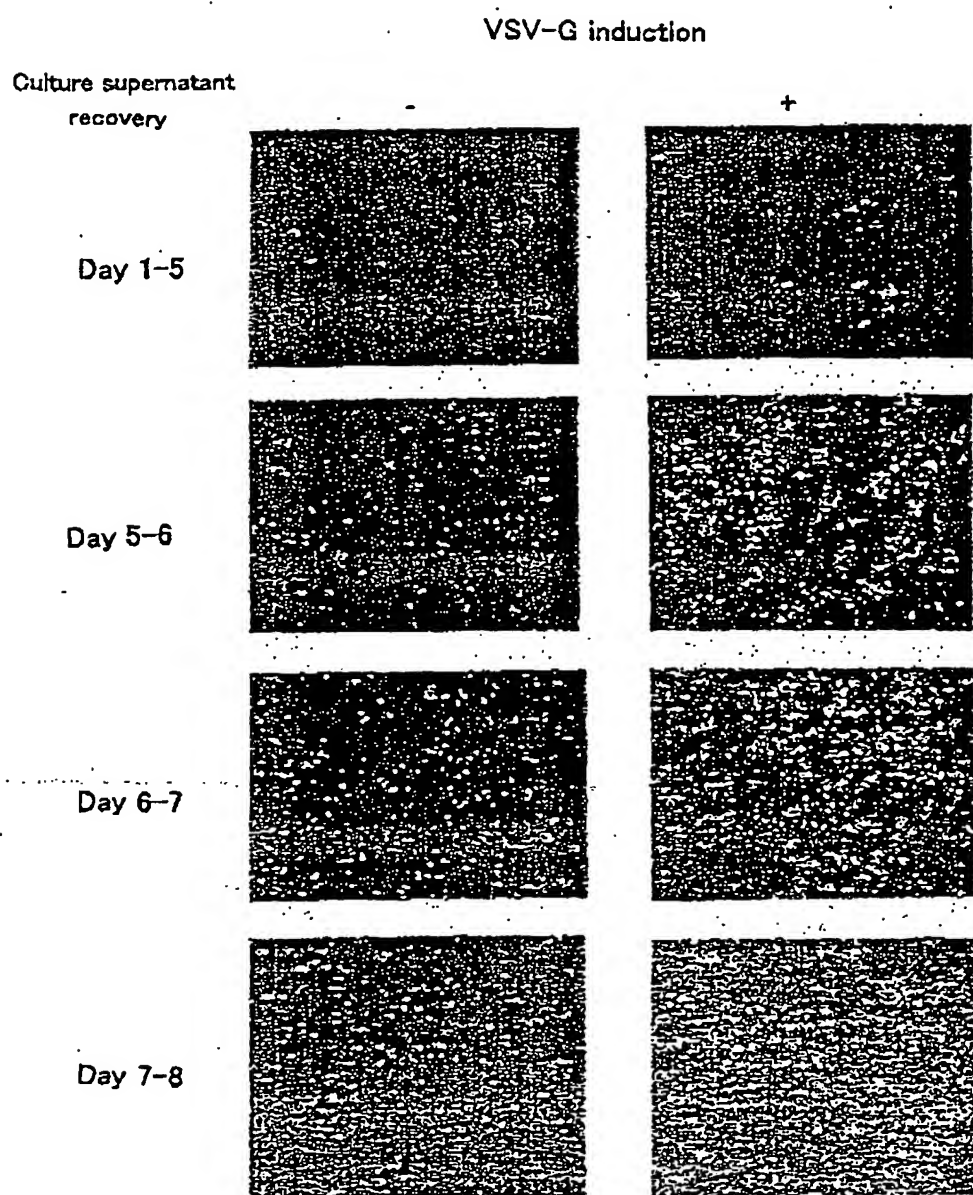


Figure 37

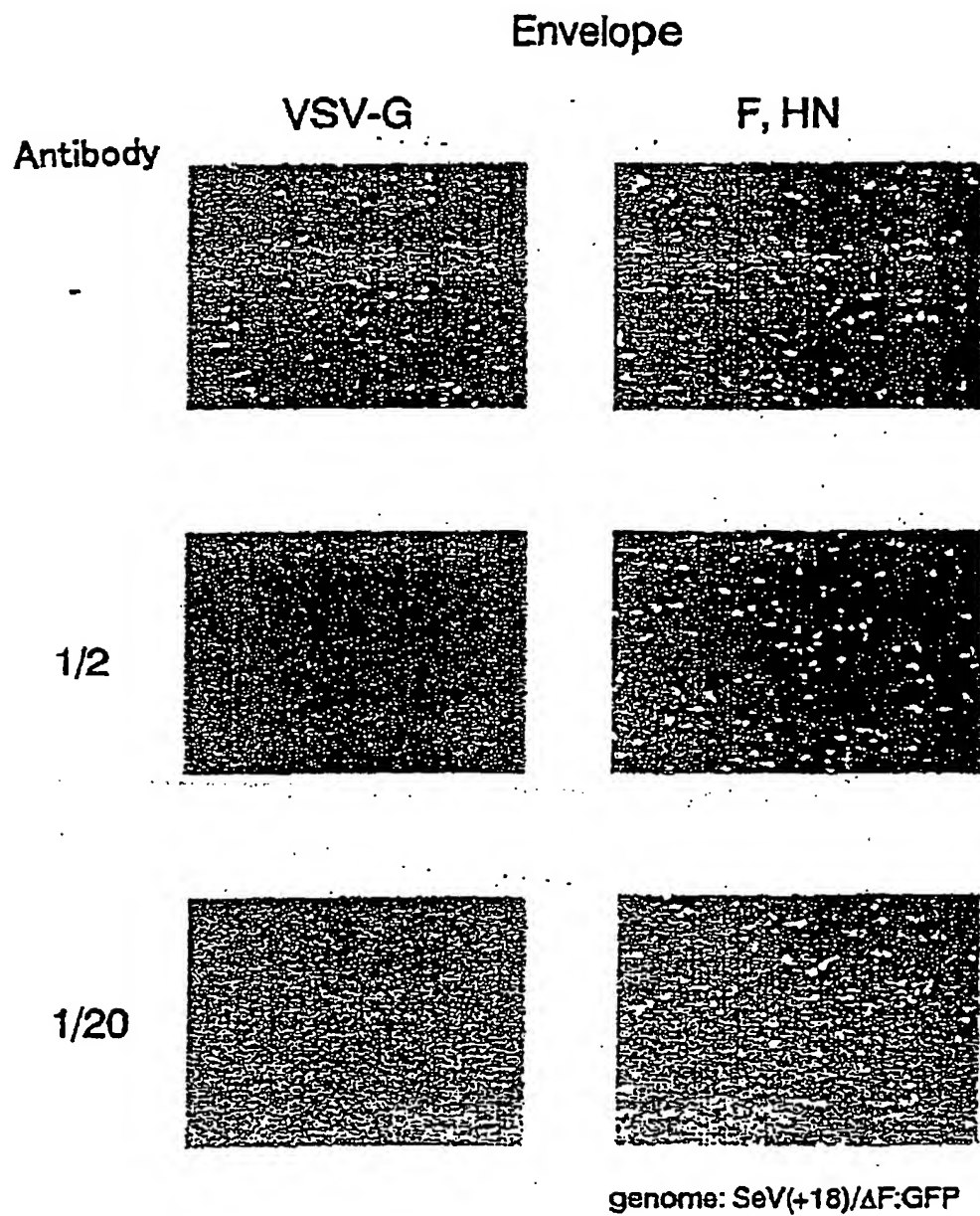


Figure 38

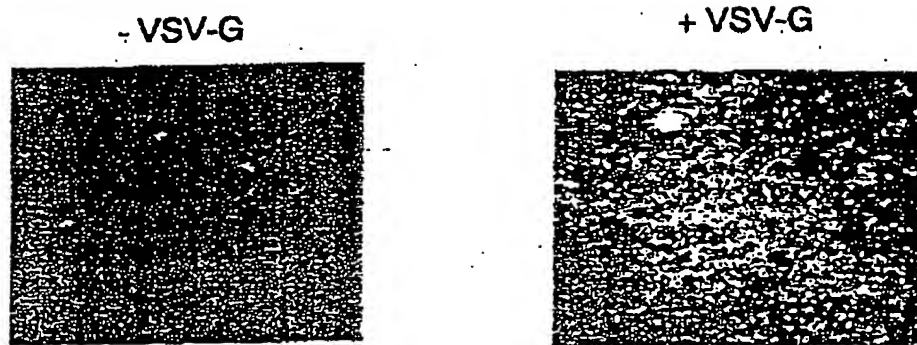


Figure 39

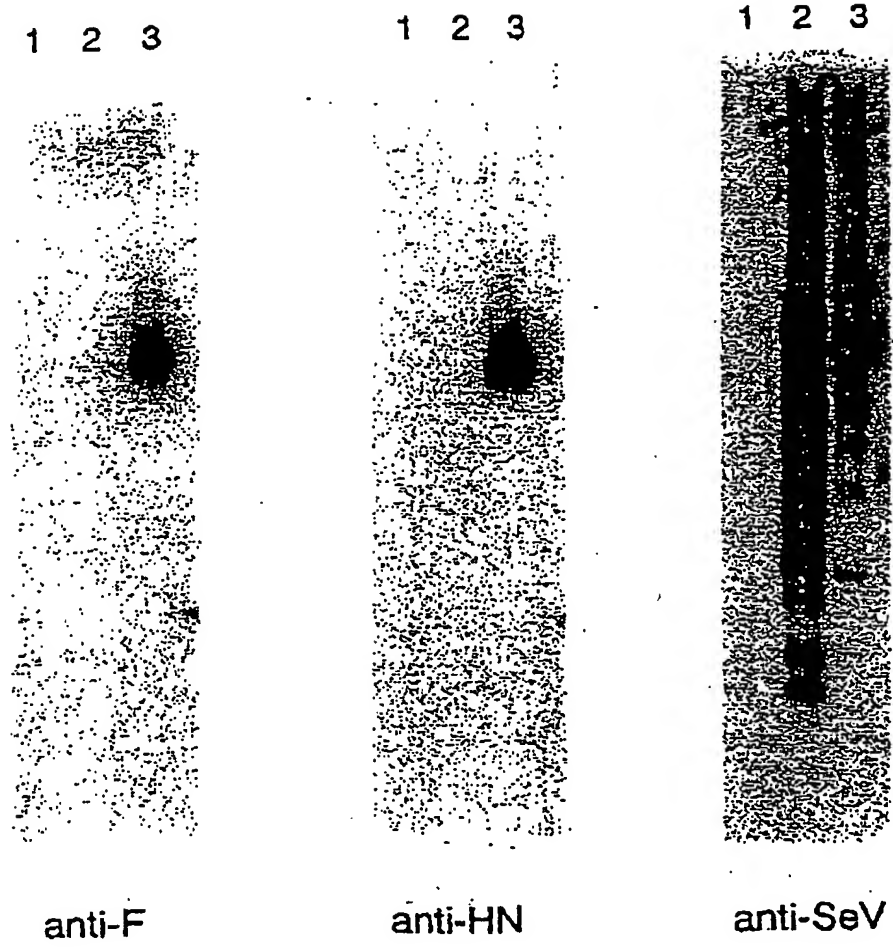
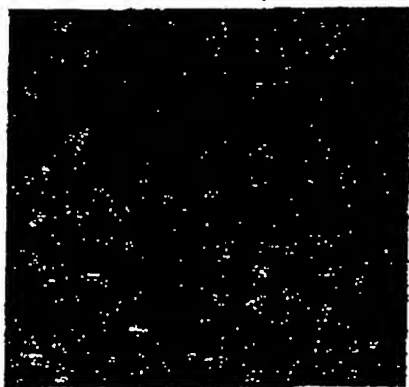
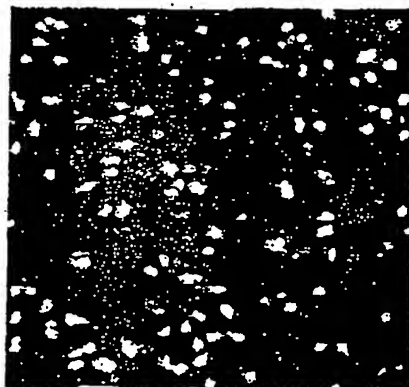


Figure 40



CIU measurement by using anti-SeV
antibody and anti-rabbit fluorescent-labeled
secondary antibody



CIU measurement of GFP-expressing cells
(control)

Figure 41

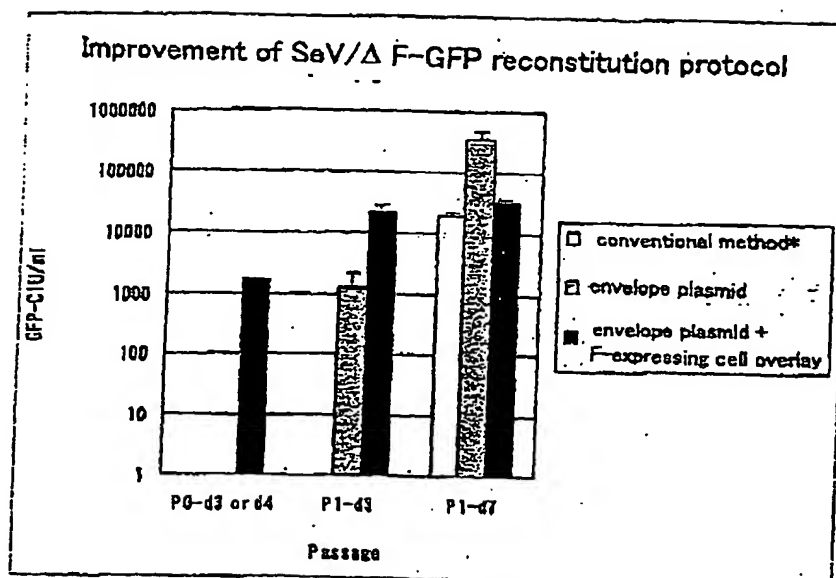


Figure 42

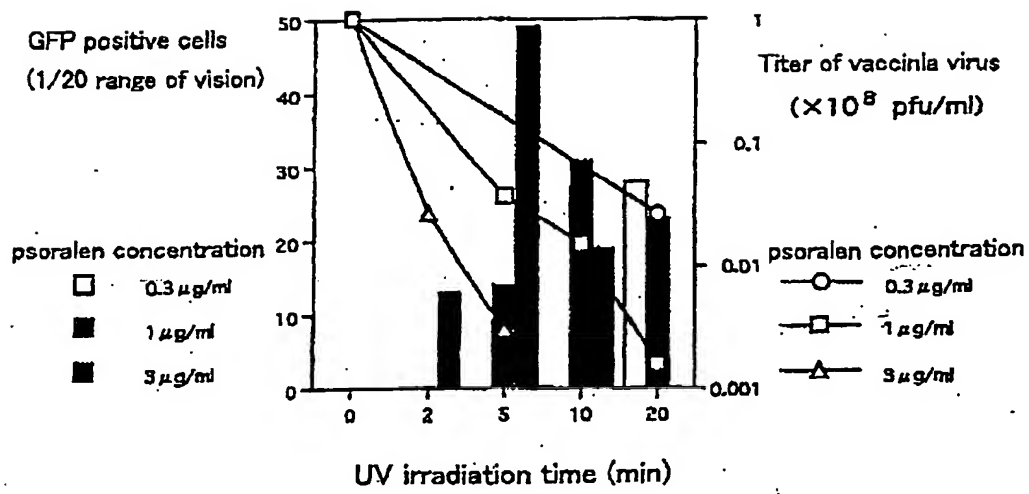
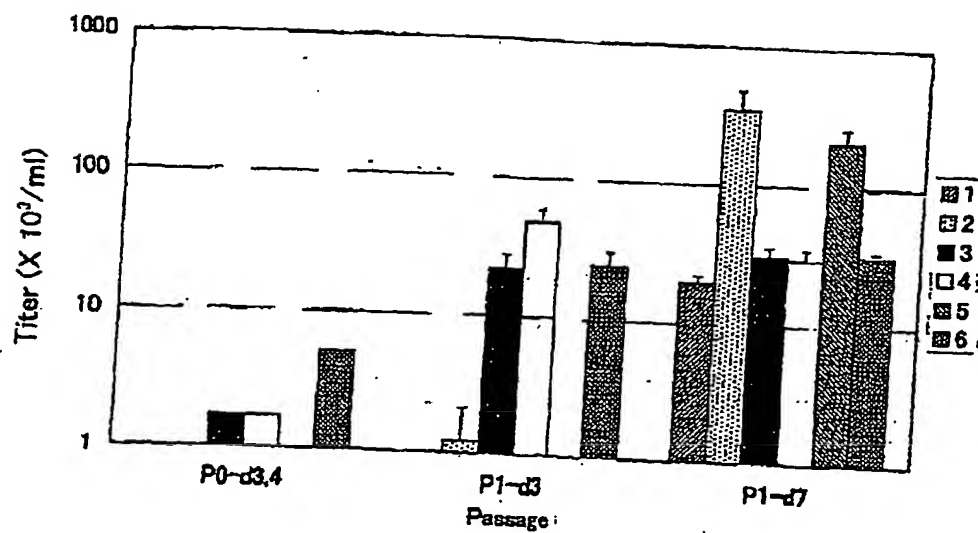


Figure 43



	V.V	V.V Lot.No	pGEM-FHN	O.L	P1-Tf.
1	VTF7-3(3,5) ϕ 35mm*	990913	-	-	Dosper
2	VTF7-3(3,5) ϕ 35mm*	990913	○	○	O.L
3	VTF7-3(3,5) ϕ 35mm*	990913	○	○	O.L
4	VTF7-3(3,5) ϕ 35mm*	000202	○	○	O.L
5	VTF7-3(0,0)	990913	○	○	O.L
6	VTF7-3(3,5) ϕ 100mm*	000202	○	○	O.L

* : Dish size used for PLWUV treatment

in O : (psoralen concentration μ g/ml, UV time, min)

Figure 44

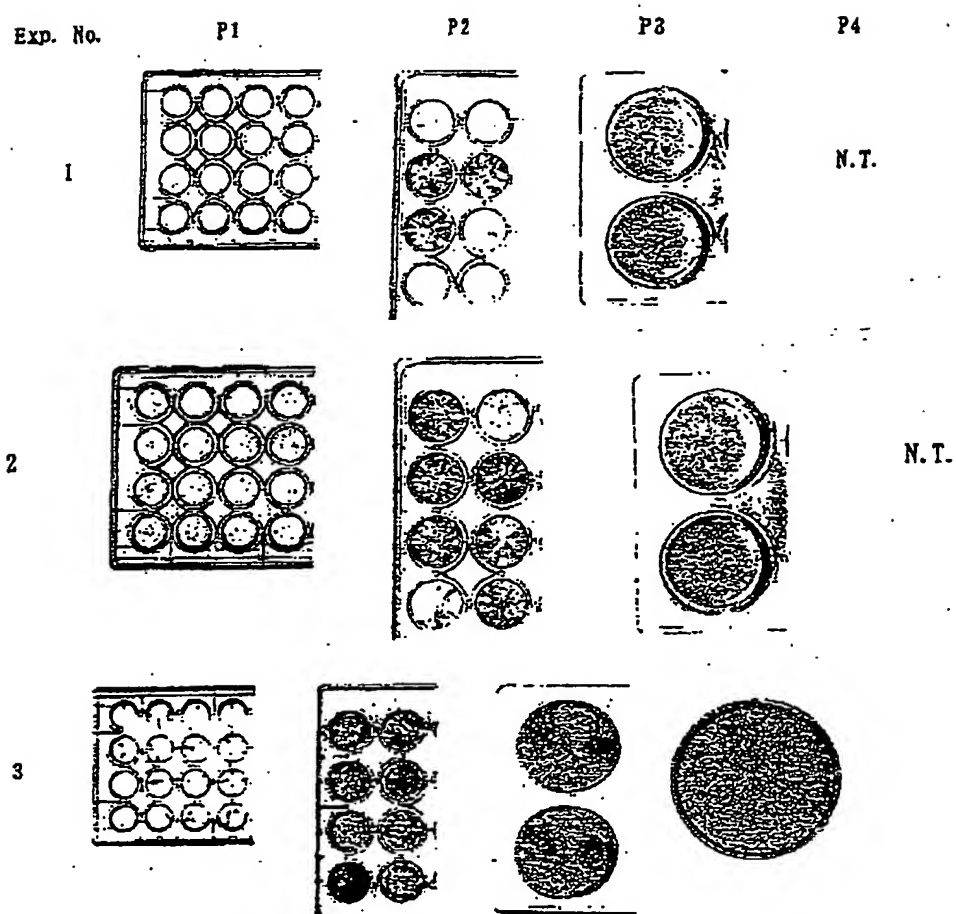
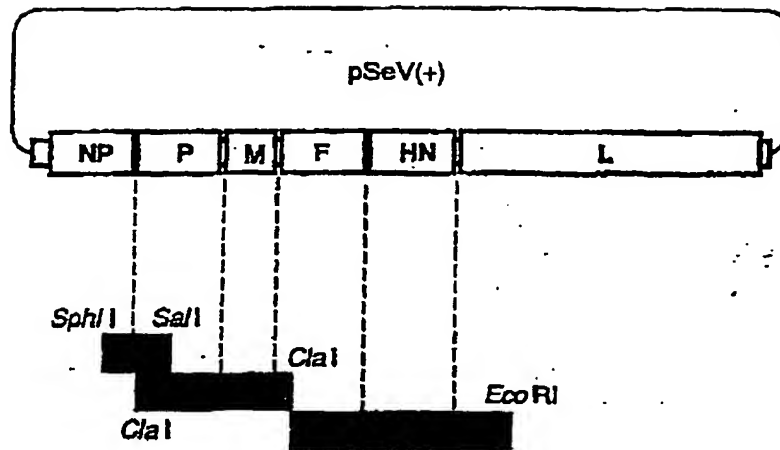


Figure 45

(A)



(B)

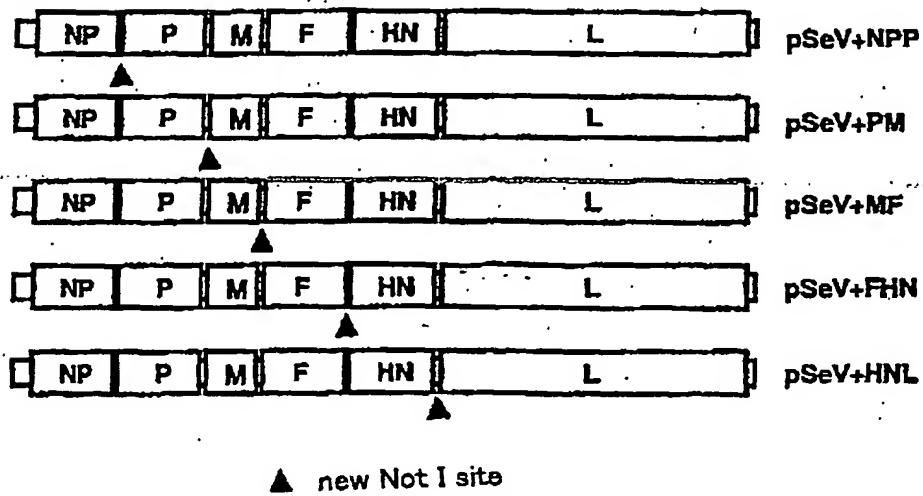


Figure 46

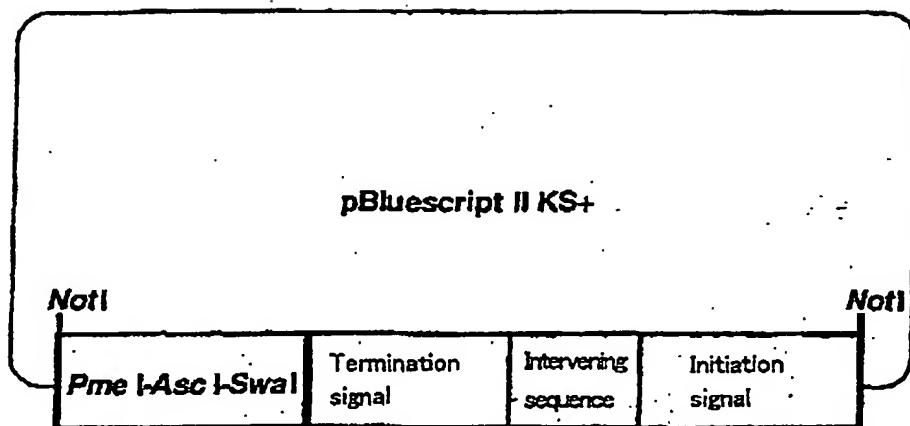


Figure 47

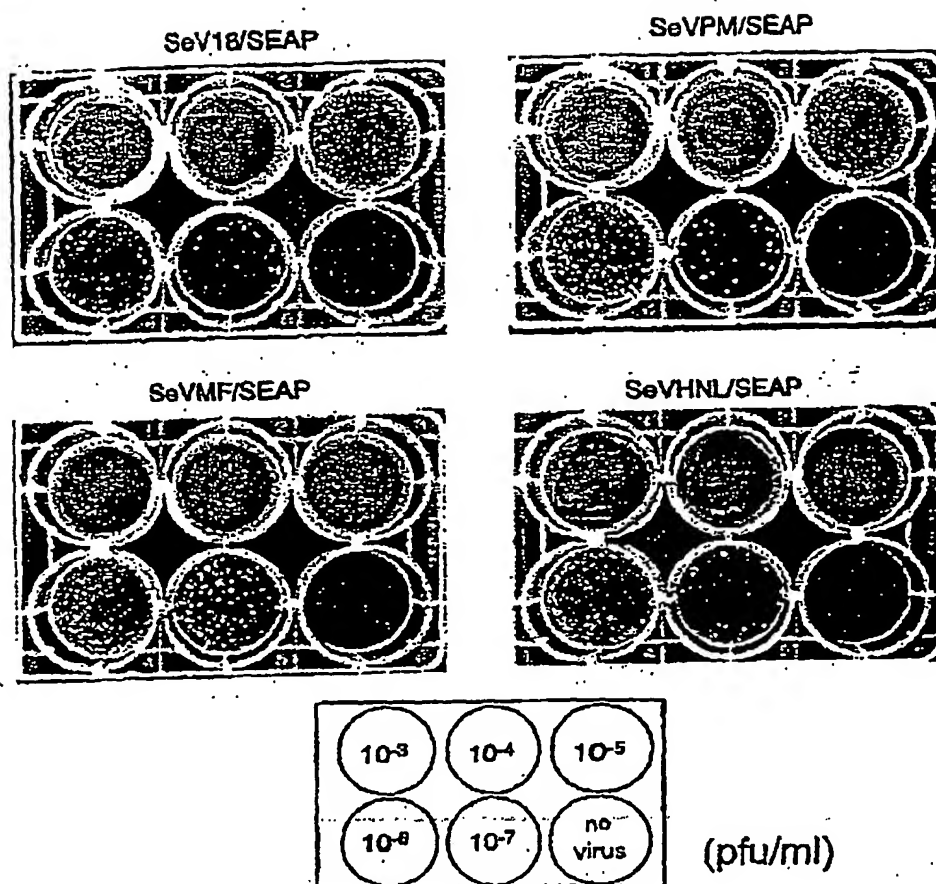


Figure 48

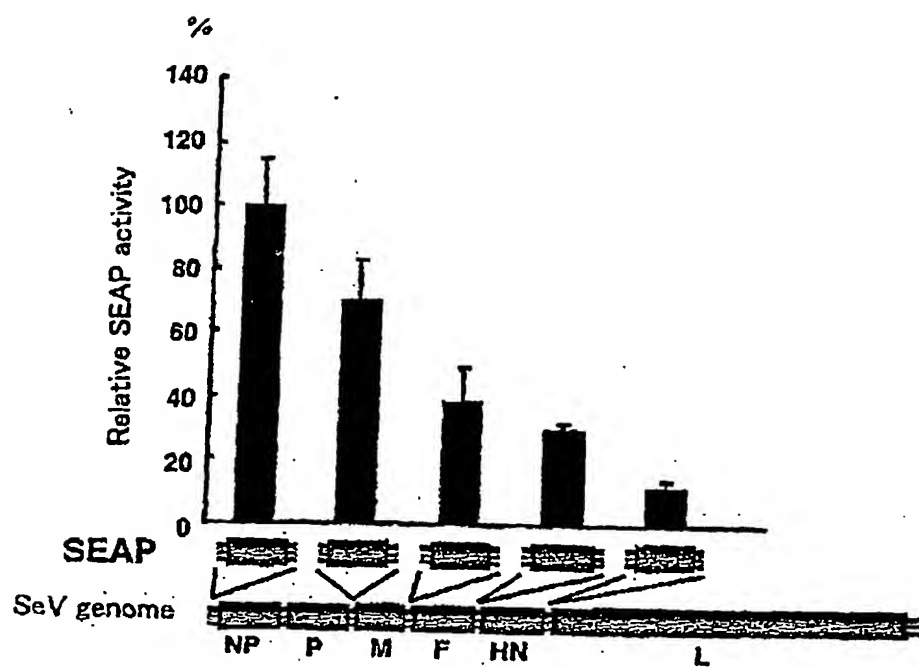


Figure 49

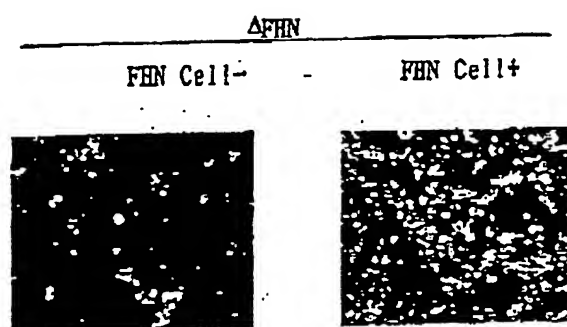


Figure 50

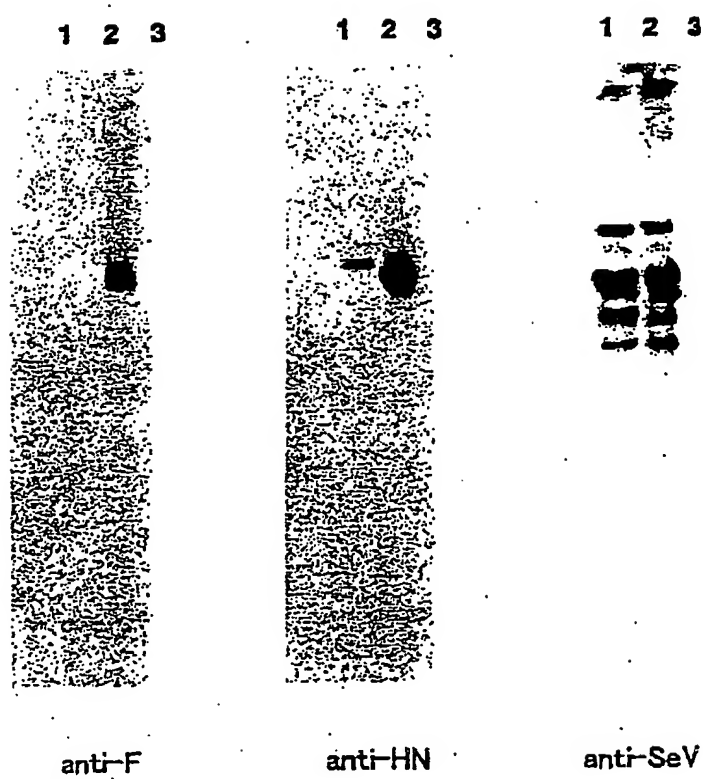


Figure 51

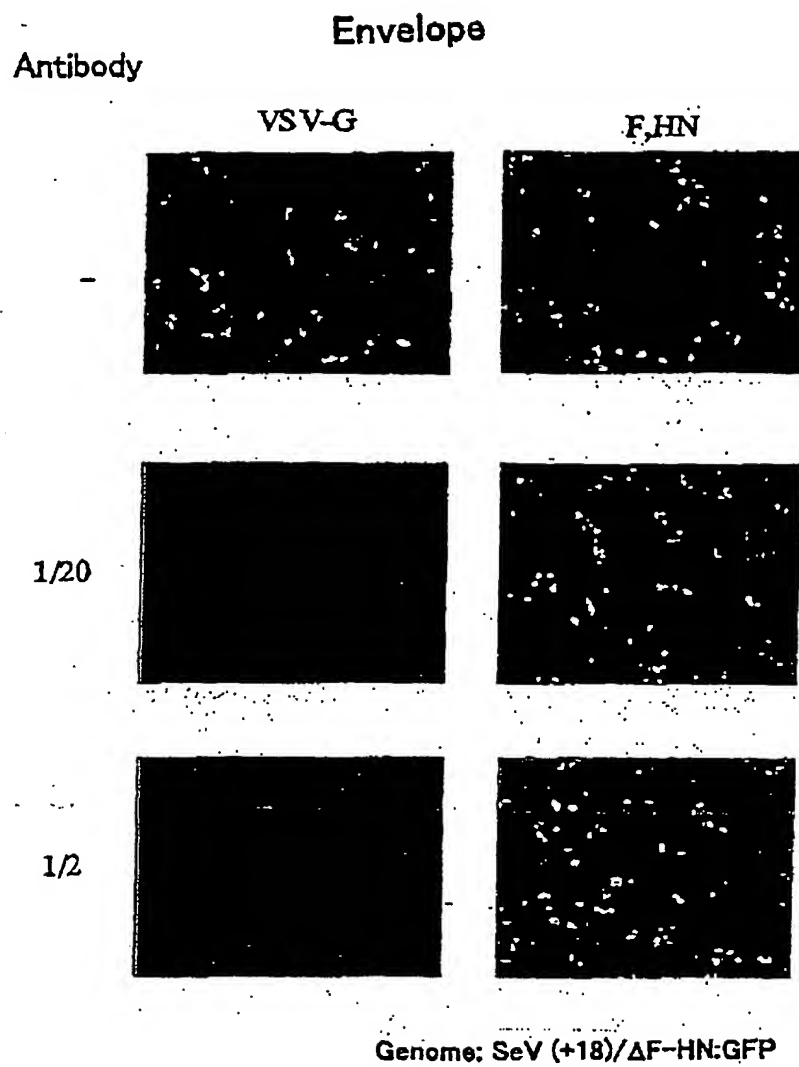
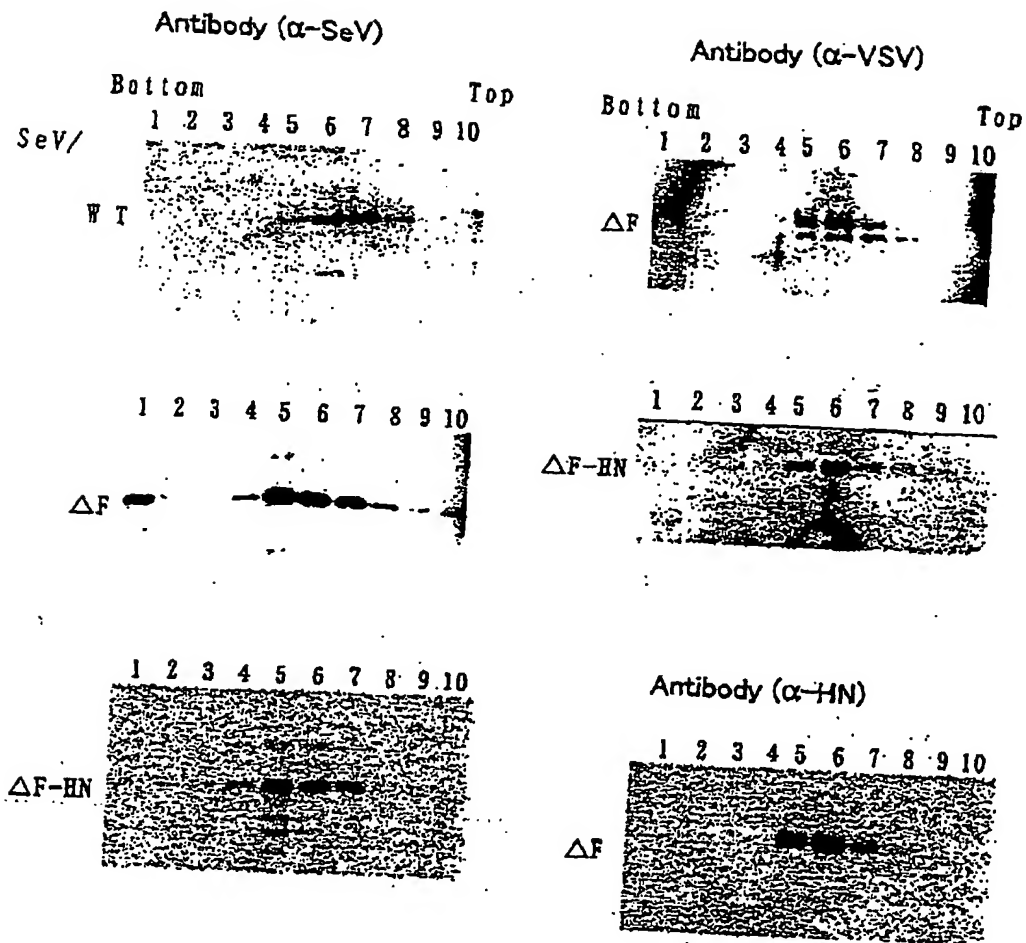
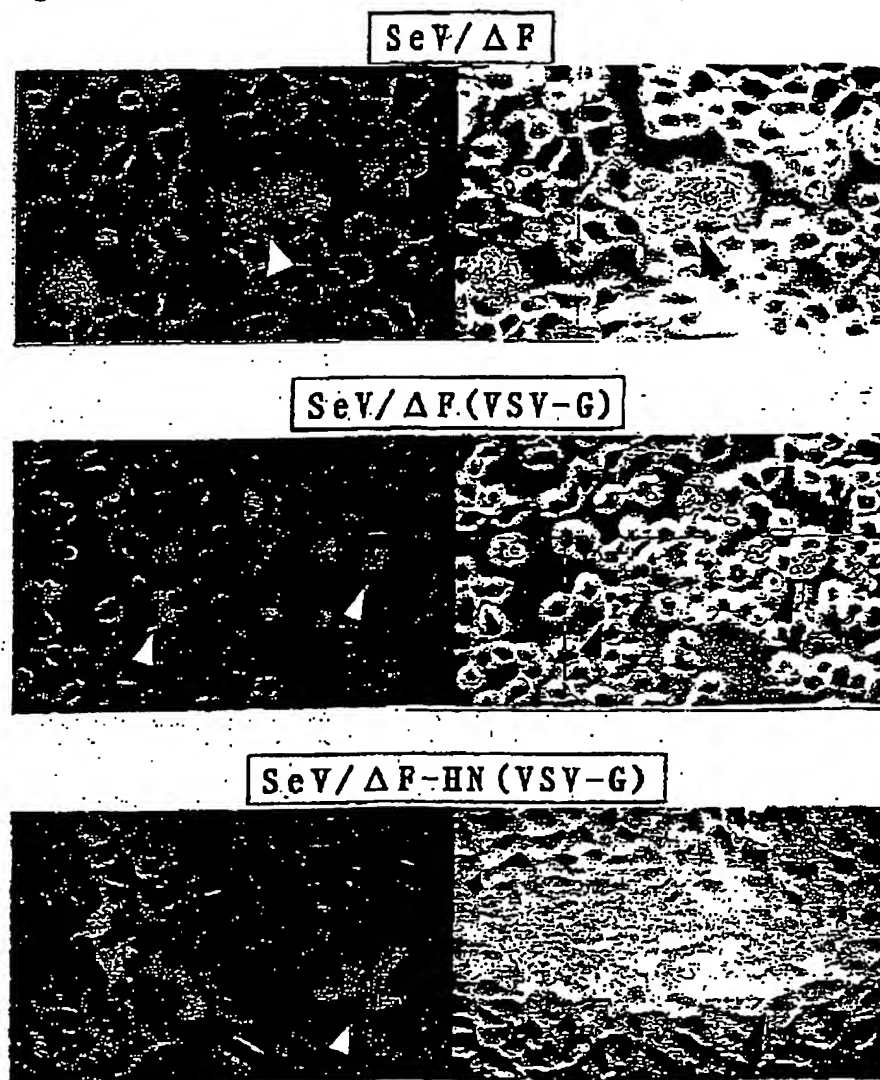


Figure 52



Sucrose: 20 - 60% linear gradient

Figure 53



(LLC-MK2 cells)

Figure 54

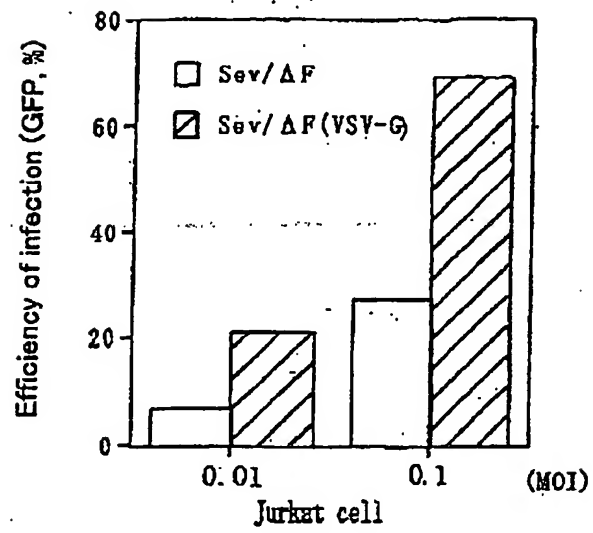
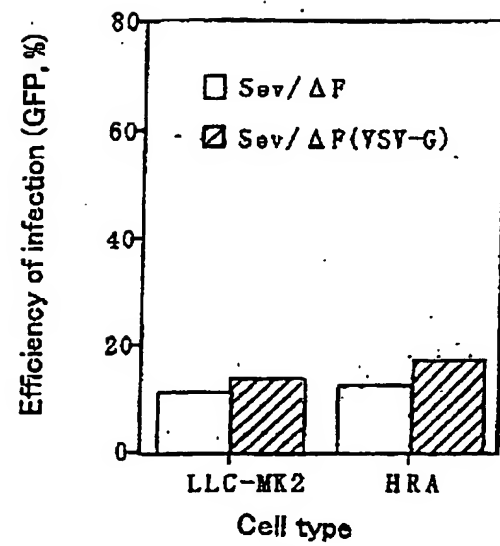
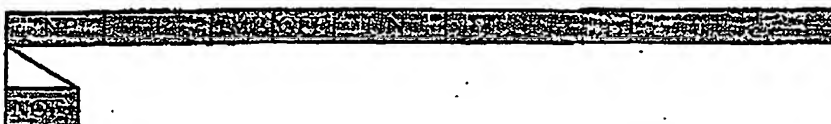


Figure 55

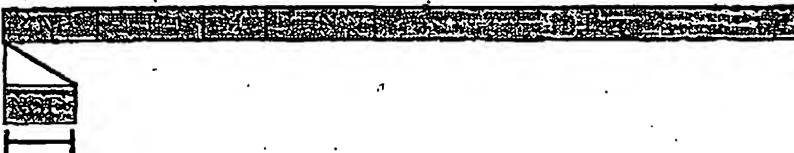
pSeV18⁺b(+)



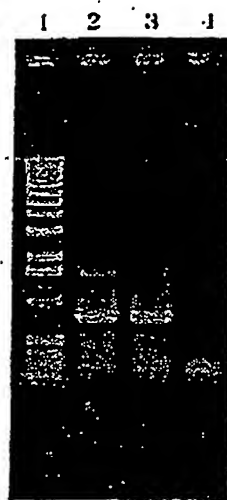
NGF/SeV/ΔF-GFP



NGF/SeV/ΔF



NGF primer (NGF-N to NGF-C)



1. Marker
2. NGF/SeV/ΔF
3. NGF/SeV/ΔF-GFP
4. pSeV18⁺b(+)

Figure 56

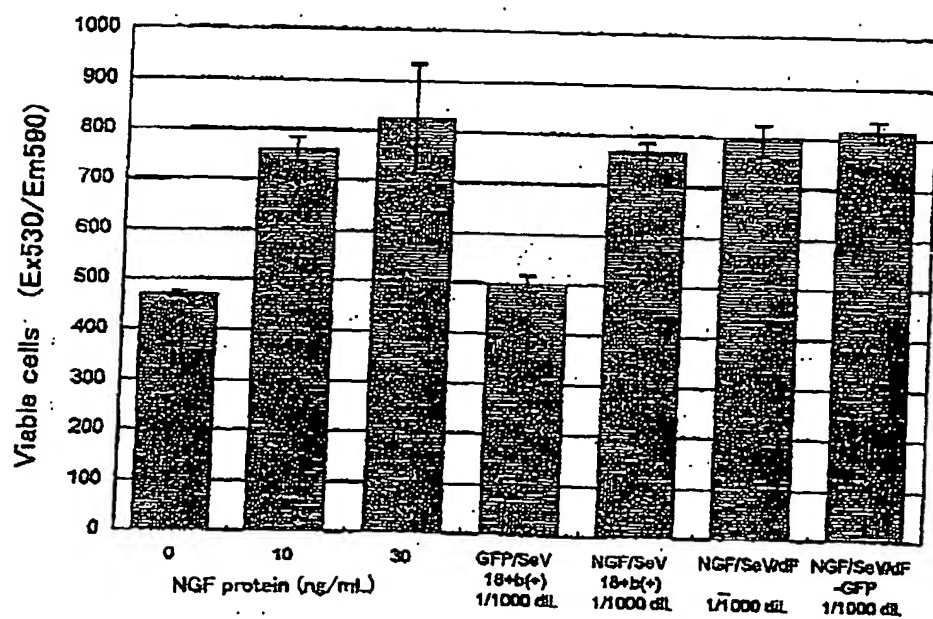


Figure 57

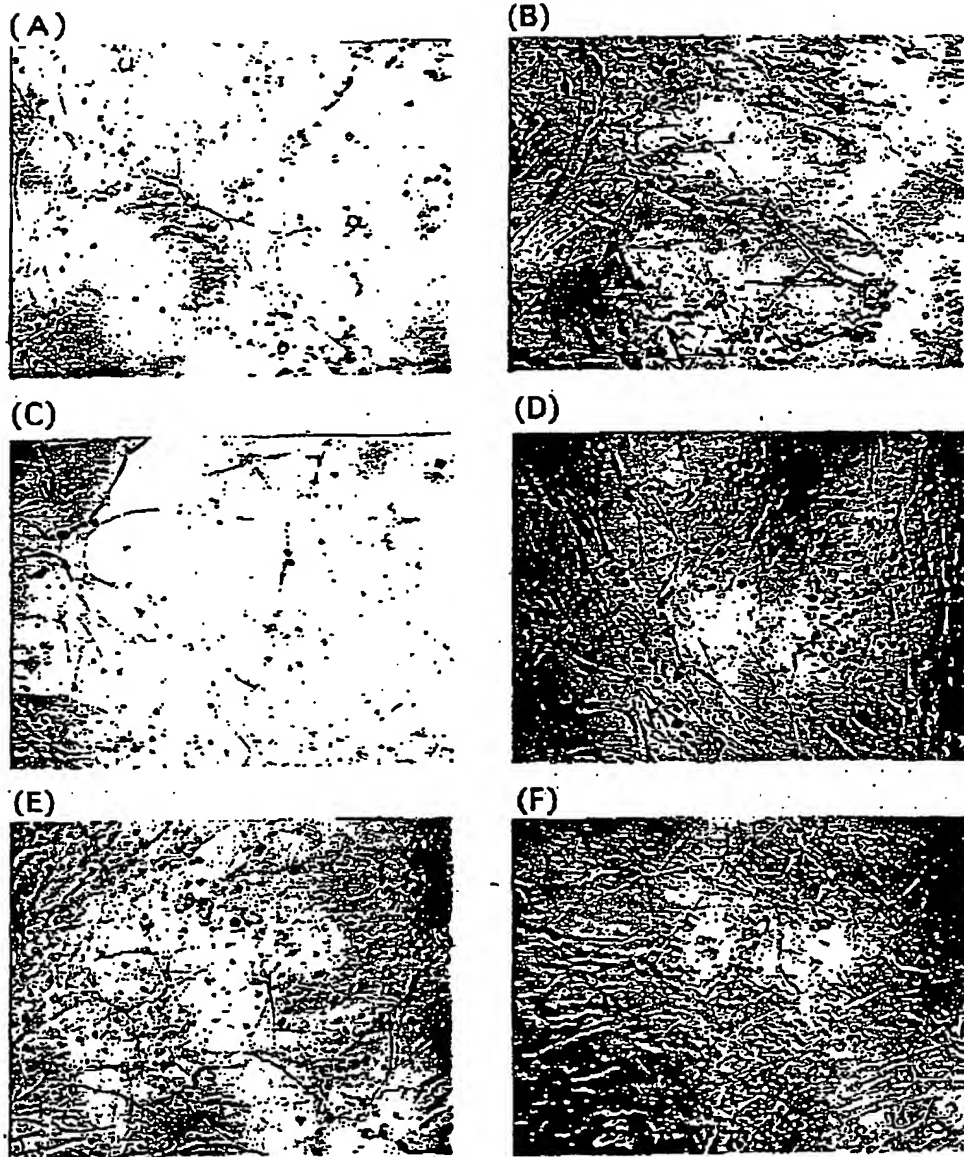


Figure 58

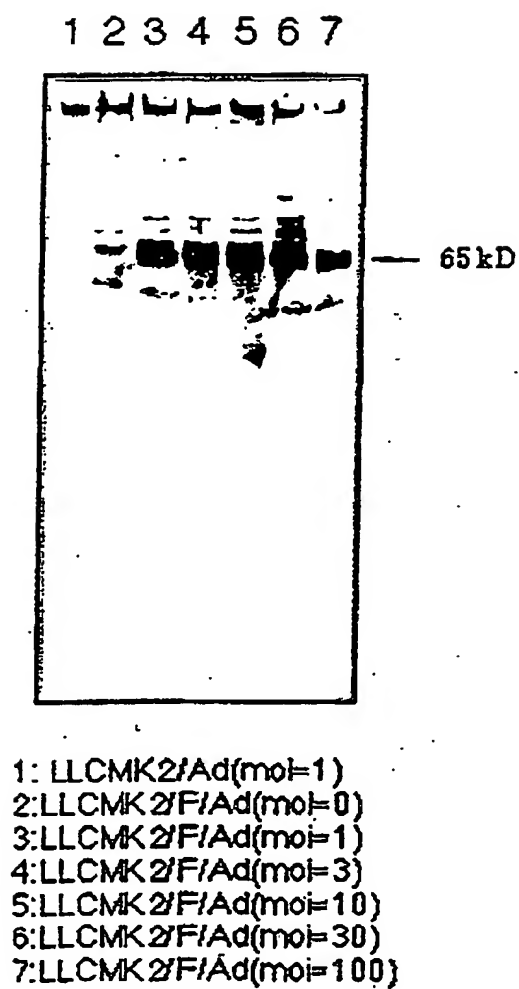


Figure 59

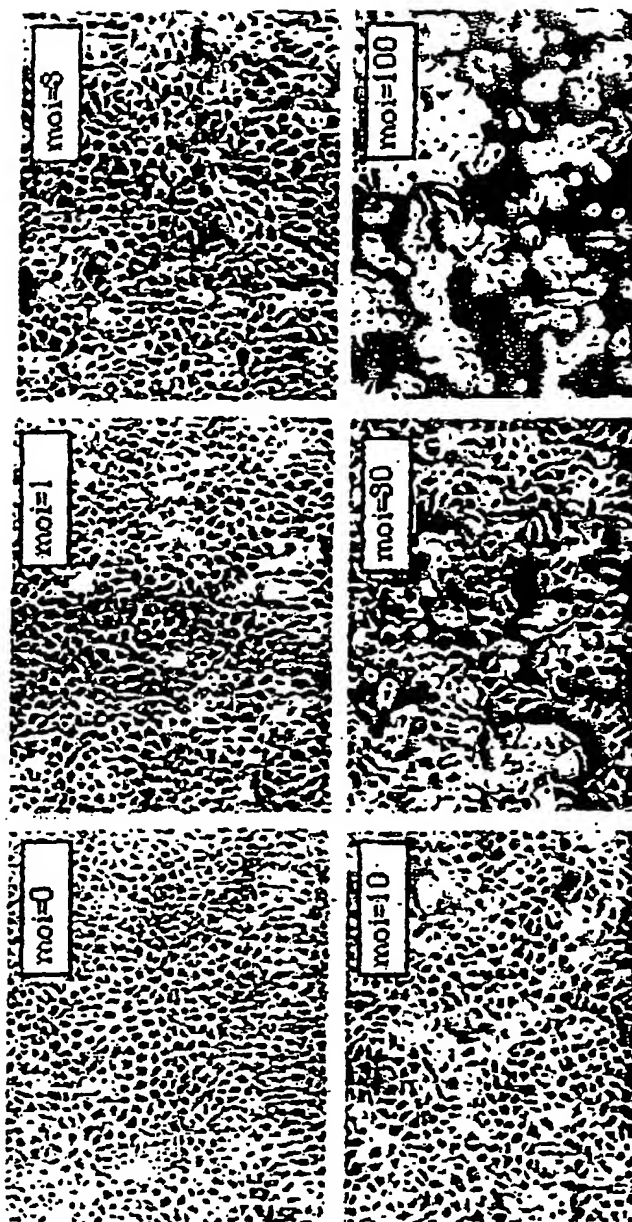
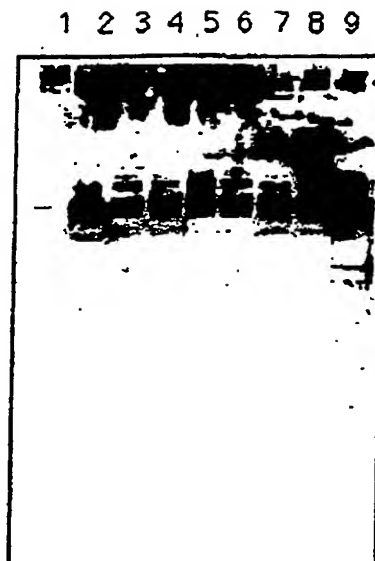


Figure 60



- 1 : LLC-MK2/Ad-Cre(Negative Control)
- 2 : LLC-MK2/F/Ad-Cre /P0
- 3 : LLC-MK2/F/Ad-Cre /P1
- 4 : LLC-MK2/F/Ad-Cre /P2
- 5 : LLC-MK2/F/Ad-Cre /P3
- 6 : LLC-MK2/F/Ad-Cre /P4
- 7 : LLC-MK2/F/Ad-Cre /P5
- 8 : LLC-MK2/F/Ad-Cre /P6
- 9 : LLC-MK2/F/Ad-Cre /P7

Figure 61

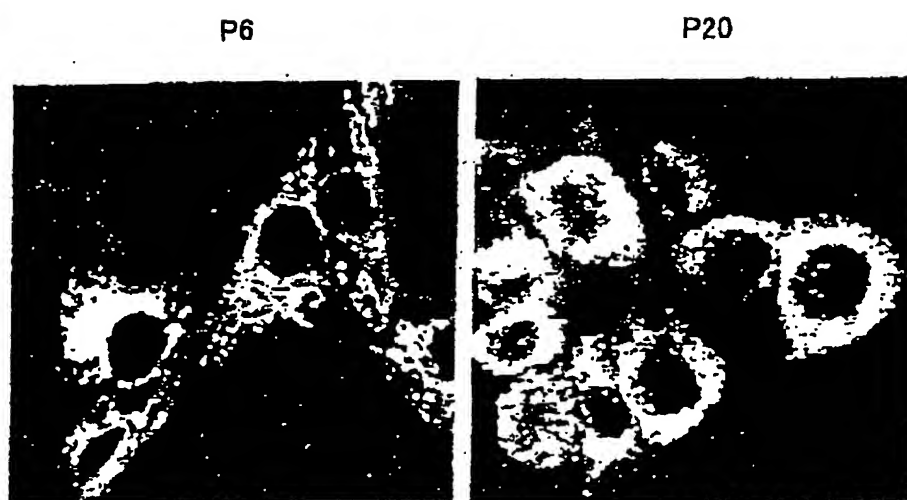
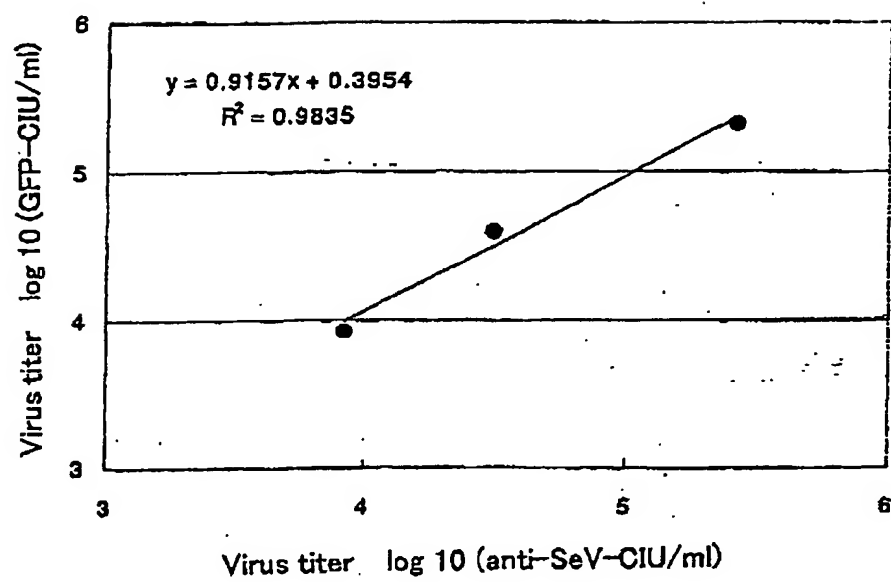


Figure 62



INTERNATIONAL SEARCH REPORT

International application No.

PCT/JP00/03194

A. CLASSIFICATION OF SUBJECT MATTER
Int.Cl.⁷ C12N15/45, C12P21/02

According to International Patent Classification (IPC) or to both national classification and IPC

B. FIELDS SEARCHED

Minimum documentation searched (classification system followed by classification symbols)

Int.Cl.⁷ C12N15/45, C12P21/02

Documentation searched other than minimum documentation to the extent that such documents are included in the fields searched

Electronic data base consulted during the international search (name of data base and, where practicable, search terms used)
WPI, WPI/L, BIOSIS PREVIEWS, CAS ONLINE, GenBank/EMBL/DDBJ/Geneseq

C. DOCUMENTS CONSIDERED TO BE RELEVANT

Category*	Citation of document, with indication, where appropriate, of the relevant passages	Relevant to claim No.
A	Huntley CC et al., "Phosphorylation of Sendai virus phosphoprotein by cellular protein kinase C zeta.", J. Biol. Chem. (1997), Vol.272, No.26, pp.16578-16584	1-7
A	Rhironov OP et al., "Solubilization of matrix protein M1-M from virions occurs at different pH for orthomyxoviruses and paramyxoviruses", Virology (1990), Vol.176, No.1, pp.274-279	1-7

☐ Further documents are listed in the continuation of Box C.☐ See patent family annex.

* Special categories of cited documents:

"A" document defining the general state of the art which is not considered to be of particular relevance

"E" earlier document but published on or after the international filing date

"L" document which may throw doubts on priority claim(s) or which is cited to establish the publication date of another citation or other special reason (as specified)

"O" document referring to an oral disclosure, use, exhibition or other means

"P" document published prior to the international filing date but later than the priority date claimed

"T" later document published after the international filing date or priority date and not in conflict with the application but cited to understand the principle or theory underlying the invention

"X" document of particular relevance; the claimed invention cannot be considered novel or cannot be considered to involve an inventive step when the document is taken alone

"Y" document of particular relevance; the claimed invention cannot be considered to involve an inventive step when the document is combined with one or more other such documents, such combination being obvious to a person skilled in the art

"&" document member of the same patent family

Date of the actual completion of the international search
15 August, 2000 (15.08.00)

Date of mailing of the international search report
12 September 2000 (12.09.00)

Name and mailing address of the ISA/
Japanese Patent Office

Authorized officer

Facsimile No.

Telephone No.

Form PCT/ISA/210 (second sheet) (July 1992)

**This Page is Inserted by IFW Indexing and Scanning
Operations and is not part of the Official Record**

BEST AVAILABLE IMAGES

Defective images within this document are accurate representations of the original documents submitted by the applicant.

Defects in the images include but are not limited to the items checked:

- ☐ BLACK BORDERS
- ☐ IMAGE CUT OFF AT TOP, BOTTOM OR SIDES
- ☐ FADED TEXT OR DRAWING
- ☒ BLURRED OR ILLEGIBLE TEXT OR DRAWING
- ☐ SKEWED/SLANTED IMAGES
- ☐ COLOR OR BLACK AND WHITE PHOTOGRAPHS
- ☐ GRAY SCALE DOCUMENTS
- ☐ LINES OR MARKS ON ORIGINAL DOCUMENT
- ☐ REFERENCE(S) OR EXHIBIT(S) SUBMITTED ARE POOR QUALITY
- ☐ OTHER: _____

IMAGES ARE BEST AVAILABLE COPY.

As rescanning these documents will not correct the image problems checked, please do not report these problems to the IFW Image Problem Mailbox.